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1 Degradation of a novel DNA damage response protein, tankyrase 1 binding protein 1 (Tab182),
2 following adenovirus infection.
3
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25 **Abstract**

26 Infection by most DNA viruses activates a cellular DNA damage response (DDR), which may be to the
27 detriment or advantage of the virus. In the case of adenoviruses, they neutralise anti-viral effects of
28 DDR activation by targeting a number of proteins for rapid proteasome-mediated degradation. We
29 have now identified a novel DDR protein, tankyrase 1 binding protein 1 (TNKS1BP1 also known as
30 Tab182), which is degraded during infection by adenovirus 5 and adenovirus 12. In both cases,
31 degradation requires the action of E1B55K and E4orf6 viral proteins and is mediated through the
32 proteasome by the action of cullin-based cellular E3 ligases. The degradation of Tab182 appears to
33 be serotype specific as the protein remains relatively stable following infection with adenoviruses 4,
34 7, 9 and 11. We have gone on to confirm that Tab182 is an integral component of the CNOT
35 complex, which has transcriptional regulatory, deadenylation and E3 ligase activity. At least 2 other
36 members of the complex (CNOT3 and CNOT7) are also reduced in level during adenovirus infection
37 whereas levels of CNOT4 and CNOT1 remain stable. Depletion of Tab182 with siRNA enhances
38 expression of E1As to a limited extent during adenovirus infection but depletion of CNOT1 is
39 particularly advantageous to the virus and results in a marked increase in expression of adenovirus
40 early proteins. In addition, depletion of Tab182 and CNOT1 results in a limited increase in viral DNA
41 during infection. We conclude that the cellular CNOT complex is a previously unidentified major
42 target for adenoviruses during infection.

43 **Importance**

44 Adenoviruses target a number of cellular proteins involved in the DNA damage response for rapid
45 degradation. We have now shown that Tab182, which we have confirmed to be an integral
46 component of the mammalian CNOT complex, is degraded following infection by adenovirus
47 serotypes 5 and 12. This requires the viral E1B55K and E4orf6 proteins and is mediated by cullin-
48 based E3 ligases and the proteasome. In addition to Tab182, other CNOT proteins are also reduced
49 during adenovirus infection. Thus, CNOT3 and CNOT7, for example, are degraded whereas CNOT4
50 and CNOT1 are not. siRNA-mediated depletion of components of the complex enhances the

51 expression of adenovirus early proteins and increases the concentration of viral DNA produced
52 during infection. This study highlights a novel protein complex, CNOT, which is targeted for
53 adenovirus-mediated protein degradation. To our knowledge this is the first time that the CNOT
54 complex has been identified as an adenoviral target.

55 **Introduction**

56 Adenoviruses are, together with the Papilloma and Polyoma viruses, members of the small DNA
57 tumour virus family (1). There are in excess of 70 human adenovirus types, subdivided into 7 species,
58 designated A-G; the most commonly studied are the group C adenoviruses types 2 and 5 (Ad2 and
59 Ad5) and the group A oncogenic adenovirus 12 (Ad12). Adenoviruses have a linear double-stranded
60 DNA genome, approximately 35kbp in length. The first gene to be expressed, following infection, is
61 adenovirus early region 1A (AdE1A) which is present in two major forms-a long form and a short
62 form translated from 13S and 12S mRNAs, respectively. AdE1A induces progression of the host cell
63 into a 'pseudo-S-phase' through interaction with a number of cellular proteins, such as the Rb family,
64 CBP/p300 and components of the cellular transcriptional machinery (2-4). It is considered that this
65 provides an environment conducive to viral replication. Adenovirus E1A is the major adenovirus
66 oncogene and has long been known to transform cells in culture in combination with a co-operating
67 oncogene, such mutant Ras or adenovirus E1B (3, 5).

68 Shortly after initial infection the host cell initiates a DNA damage response (DDR), seen as
69 phosphorylation of a number of well-characterised Ataxia telangiectasia mutated (ATM) and ATM
70 and Rad3-related (ATR) substrates (6-8). It is presumed that this may be due to recognition of the
71 viral genome as broken cellular DNA or perhaps due to stress caused by infection itself. The virus, in
72 turn, is able to inhibit the DDR, primarily by degradation and/or mis-localization of its key
73 components (7-12). The cellular DDR comprises a series of pathways which have evolved to deal
74 with different forms of DNA damage, such as double strand breaks (DSBs), single strand breaks
75 (SSBs), and the formation of bulky adducts and base mismatches (13-15). The response to DSBs is
76 largely based on the activities of three kinases – ATM, ATR, and DNA dependent protein kinase

(DNA-PK). DSBs can be detected by both the MRN complex (comprising MRE11, Rad50 and NBS1) or the Ku 70/80 heterodimer which can lead to repair by homologous recombination (HR) or non-homologous end joining (NHEJ), respectively. Recognition of DSBs by MRN is followed by the recruitment of ATM, which is activated by acetylation by Tip60 while binding of Ku70/80 results in auto-phosphorylation of DNA-PK that is required for NHEJ (16-18). Histone H2AX and multiple downstream targets are phosphorylated by ATM which has the effect of recruiting a large number of components to the lesion to initiate repair, as well as to cause cell cycle arrest so that damaged DNA is not replicated (13-17). ATR is activated in response to single-stranded DNA (ssDNA) which can arise as a result of DSB repair and stalled replication forks. Regions of ssDNA are coated with replication protein A (RPA) which, in turn, recruits ATR and the ATR interacting protein, ATRIP. Further complexes, comprising Rad9-Rad1 and Hus1 (9-1-1) and Rad17-replication factor C2 (RFC2) clamp loader, together with TOPBP1 are recruited to ssDNA, RPA and ATR leading to cell cycle arrest and repair (17-19).

It was originally shown that when cells were infected with a mutant Ad5 virus, lacking the E4 region, viral genomes were joined end to end to form concatamers which could not be packaged into viral capsids (20). It was later demonstrated that during infection with *wt* virus, cellular E3 ligases are hijacked by the virus and used to degrade key cellular DDR proteins; for example, p53 is degraded by both Ad5 and Ad12 and requires the action of the E1B55K and E4orf6 viral proteins (21-23). In the case of Ad5 the viral proteins recruit an E3 ligase, comprising elongins B and C, Rbx and cullin 5, which ubiquitylates p53 and this is then degraded by the proteasome (9). Similarly, Ad12 also facilitates the degradation of p53, but through a cullin 2-based E3 ligase (12). Other DDR proteins degraded during Ad5 and Ad12 infection include MRE11, DNA ligase IV and BLM (10, 11, and 24). In addition to DDR components a number of other unrelated proteins are also degraded during Ad5 infection. These substrates include DAXX, integrin3 α and TIF1 γ (25-27). During infection adenoviruses also cause translocation of proteins associated with the DDR. For example, ATR, ATRIP, Rad 17, 53BP1, BRCA1, TOPBP1, RPA and hnRNPUL-1 have all been observed at sites of viral

103 replication in the nucleus, known as viral replication centres (VRCs) (6, 8 and 28). In addition, it is
104 notable that certain DDR proteins, such as p53 and MRE11, are translocated to aggresomes where
105 they may be degraded (29-31).

106 Tab182 (also known as tankyrase 1 binding protein 1 [TNKS1BP1]) has previously been shown to be
107 an ATM and/or ATR substrate which is highly phosphorylated following exposure to ionizing
108 radiation (IR) and to bind to tankyrase 1 (32, 33). It appears to be required for efficient DSB repair
109 and facilitates PARP1-dependent autophosphorylation of DNA-PK although its precise role is not
110 clear at present (34, 35 and our unpublished data). In addition, Tab182 has a role in regulation of the
111 actin cytoskeleton (36). Tab182 has previously been suggested to be a component of the mammalian
112 CNOT complex although its role in that context is unknown. The CNOT complex is a multi-protein
113 complex, highly conserved in eukaryotes (37-39). In humans, the CNOT complex is composed of
114 components CNOT1 to 11 (CNOT9 and CNOT11 have the alternative nomenclature RQCD and
115 C2orf29, respectively) (40, 41). In yeast, where most studies of CCR4-NOT have been performed,
116 there are 9 core subunits-Cer4, Caf1, Caf40, Caf130 and NOTs1-5, although no Tab182 ortholog has
117 been identified (38, 42 and 43). The human CNOT complex consists of a stable inner complex
118 (CNOT1, CNOT2, CNOT3, CNOT9 and CNOT10) with CNOT6 and its homologue 6L, CNOT7 and CNOT8
119 being less strongly associated. CNOT4 seems to be weakly associated, whereas Tab182 and C2orf29
120 (CNOT11) are more strongly bound (40, 44 and 45). Many different enzymatic activities have been
121 attributed to the CCR4-NOT complex in yeast and CNOT in mammals. It is considered to be a major
122 deadenylase, responsible, with Pan2-Pan3, for shortening of the poly (A) tails of cytoplasmic RNAs
123 (38, 46 and 47). The components CNOT7 and CNOT8, together with CNOT6 and CNOT6L, are
124 deadenylase subunits. Further components of the complex have E3 ligase, translational repression,
125 RNA export and nuclear surveillance activities (38, 48-50). CNOT4 is the E3 ubiquitin ligase but seems
126 to interact only weakly with the remainder of the complex (40). CNOT1 forms a scaffold on which
127 the CNOT and deadenylase modules are formed (41, 51 and 52). The central region of CNOT1
128 interacts with the deadenylase subunits, with CNOT7 forming a bridge between CNOT1 and CNOT6L

129 (39). The C-terminal region of CNOT1 binds to the remainder of the NOT module which comprises
130 CNOT2 and CNOT3.

131 A number of studies have implicated the CCR4-NOT complex in the DDR in yeast. In the majority of
132 these, sensitivity assays were performed using yeast strains that were mutant for various CCR4-NOT
133 components. For example, loss of CCR4 and Caf1 render the yeast sensitive to IR, hydroxyurea (HU)
134 and camptothecin, an inhibitor of DNA topoisomerase I (53-55). Similarly, NOT1-5 mutant yeast
135 strains have been shown to be sensitive to HU (53). These data suggest that the CCR4-NOT complex
136 is involved in the response to a number of forms of DNA damage and replication stress although the
137 mechanisms involved remain unclear.

138 Here we demonstrate that Tab182 is degraded during Ad5 and Ad12 infection in an E1B55K- and
139 E4orf6-dependent manner. We have confirmed that Tab182 is a component of the CNOT complex
140 and that levels of at least two other components of the complex are similarly reduced during
141 adenovirus infection. Significantly, depletion of Tab182 or disruption of the CNOT complex enhances
142 expression of adenovirus E1A at the transcriptional level early in infection.

143 **Results**

144 *Tab182 is degraded during adenovirus infection.*

145 It has previously been suggested that Tab182 may have a role in the DDR based on the observation
146 that the protein has multiple potential ATM/ATR phosphorylation sites (SQ/TQ) and is
147 phosphorylated following exposure to IR (33) as well as its recently proposed role in DSB repair (34,
148 35). In a screen to detect novel DDR components targeted by adenoviruses the effect of viral
149 infection on Tab182 was examined. It can be seen that, during both Ad5 and Ad12 infection of HeLa
150 cells, Tab182 levels decline rapidly after 24 hours (Figures 1A and 1B). It is particularly notable that
151 the levels of Tab182 increase in the initial stages of infection with both serotypes (Figure 1 and
152 succeeding figures). This appears to be a cell cycle effect, since, in nocodazole 'shake off'
153 experiments, Tab182 expression is greatest during S phase and mitosis and reduced in the G1 phase

154 of the cell cycle (data not shown). RT-PCR analysis has demonstrated that the increased protein
155 expression coincides with increases in Tab182 mRNA (data not shown).

156 *Tab182 degradation requires the E1B55K and E4orf6 viral proteins*

157 Multiple previous studies have demonstrated the roles of AdE1B55K and AdE4orf6 in targeting
158 cellular proteins for degradation (6-12). To determine whether these components are involved in the
159 observed reduction in level of Tab182, infection with a panel of mutant viruses was carried out.
160 Infection with the Ad5 (Ad5 *d/1520*) and Ad12 (Ad12*d/620*) E1B55K negative viruses had no effect on
161 the level of Tab182 (Figures 1C and 1D), indicating a requirement for the larger AdE1B protein for
162 degradation.

163 Following infection with various Ad5E4 negative viruses there was no reduction in Tab182 level
164 when the E4orf6 protein was not expressed, as in H5*pm4154* and H5*pm4155* (Figures 2A and 2B).
165 Viruses which fail to express other E4 proteins degrade Tab182 in a manner comparable to wild type
166 (Figure 2). Thus, H5*in351* (E4orf1-), H5*in352* (E4orf2-), H5*pm4166* (E4orf4-) and H5*pm4150* (E4orf3-)
167 are all able to cause rapid degradation of Tab182 (Figure 2). The H5*d/356* virus, which is E4orf7
168 negative, appears to express E4orf6 at much lower levels than expected which probably explains
169 why levels of Tab182 and MRE11 are only very marginally reduced (Figure 2C). H5*pm4155*, which is
170 E4orf3 and E4orf6 negative, expresses somewhat reduced levels of E1B55K, compared to the other
171 viruses shown here (Figure 2C). Reasons for this are not apparent. Overall, we conclude that
172 degradation of Tab182 requires, in Ad5 at least, E1B55K and E4orf6. Significantly, in all western blots
173 shown in Figures 1 and 2 (and Figure 5A) degradation of Tab182 occurs somewhat later than
174 degradation of MRE11 but at similar times to p53 degradation (data not shown).

175 In addition, to confirm that reduction in Tab182 levels is not due to a reduction in mRNA, RT-PCR
176 was carried out on Ad5 and Ad12 infected cells (Figure 3). This clearly shows that Tab182 mRNA
177 levels are equivalent to, or higher than, uninfected cells up to about 72 hours post infection in
178 contrast to the sharp reduction in Tab182 protein levels after 24 hours (compare Figures 1 and 3).

179 We conclude that loss of Tab182 protein is due to active protein degradation and not host cell shut-
180 off which may occur after 72 hours (Figures 3A and 3B).

181 To demonstrate that the E1B55K and E4orf6 proteins are solely responsible for degradation of
182 Tab182 plasmids encoding the two Ad5 and Ad12 proteins were transfected into HeLa cells. Cells
183 were harvested after 48 hours and lysates subjected to western blotting for Tab182, MRE11 and the
184 viral proteins (Figure 4). E4orf6 proteins were HA-tagged and were detected with an anti-HA
185 antibody. It can be seen that Tab182 and MRE11 were degraded in the presence of both Ad5 and
186 Ad12 E1B55K and E4orf6 proteins. These data confirm that similar viral proteins are required for
187 both Ad5- and Ad12-mediated degradation of Tab182. When the viral proteins were expressed
188 singly there was little reduction in Tab182 or MRE11 levels confirming that both E1B55K and E4orf6
189 are required for degradation (Figure 4).

190 *Degradation of Tab182 is limited to certain virus serotypes.*

191 To determine how widespread the degradation of Tab182 is amongst other adenovirus serotypes,
192 levels of Tab182 were monitored by western blotting following infection of HeLa cells with Ad4
193 (group E), Ad7 (group B1), Ad9 (group D) and Ad11 (group B2) (Figure 5). In contrast to Ad5 and
194 Ad12, infection of HeLa cells with Ad9 and Ad11 viruses had no effect on Tab182 expression except
195 at very late times when host cell shut off could be a contributory factor (Figures 5B and 5C).
196 Following Ad4 and Ad7 infection there is a reduction in Tab182 levels at later times and this is more
197 pronounced than the effects seen with Ad9 and Ad11 but much less marked than degradation after
198 Ad5 and Ad12 infection (Figures 5B and 5C). The effects of the viruses on Tab182 levels closely
199 mirror those on MRE11 and, in the case of Ad4, on p53 (Figure 5). (Ad7, Ad9 and Ad11 all markedly
200 induce expression of p53 as has been reported earlier (28 and 56)). We have previously reported
201 that Ad4 facilitates rapid degradation of various DDR proteins (28) although perhaps to a lesser
202 extent than Ad5 and Ad12. However, it appears to have only a relatively slight effect on Tab182
203 (Figure 5B). Whilst there is limited reduction in protein level, it seems likely that the group B1, B2, D
204 and E viruses do not cause significant degradation of Tab182.

205 *Degradation of Tab182 requires the proteasome and E3 ligases.*

206 A number of approaches were adopted to investigate the mechanism by which target proteins are
207 degraded during Ad5 and Ad12 infection. Initially, to confirm that the degradation of Tab182 is by
208 the proteasome, cells were treated with bortezomib, a well-characterised proteasome inhibitor, or
209 DMSO (as a negative control) and harvested after 48 hours. In the presence of bortezomib
210 degradation of Tab182 and MRE11, following viral infection, was reduced but not completely
211 inhibited; in the absence of the proteasome inhibitor (DMSO) the proteins were degraded in the
212 presence of the viruses (Figure 6). In a second experiment it has been shown that inhibition of
213 NEDDylation (with MLN4924) also results in stabilisation of Tab182 following Ad5 and Ad12
214 infection. It is now well-established that NEDDylation is required for activation of the cullin
215 components of the E3 ligases during adenovirus infection (9). In the presence of the inhibitor,
216 degradation of Tab182 was appreciably reduced as was that of p53, although it is interesting to note
217 that stabilization of MRE11 was appreciably less than was the case for p53; this apparent differential
218 may be due to the up-regulation of p53 expression due to AdE1A (Figure 7A and 7B). The active
219 NEDDylated component of the cullin 2 can be seen as a slower migrating protein in the western blots
220 shown in Figures 7A and 7B. This is markedly reduced in the MLN4924 treated samples. We conclude
221 that active (NEDDylated) cullins are required for Tab182 degradation during adenovirus infection.
222 Different adenovirus serotypes do not all make use of the same cullin components to degrade
223 cellular proteins. Previously it has been shown that protein degradation following Ad5 infection
224 utilizes a cullin 5-based E3 ligase whereas Ad12 hijacks a cullin 2-based E3 ligase (9 and 12). To
225 examine whether this difference extends to the degradation of Tab182, H1299 cells, in which Cul2 or
226 Cul5 expression had been ablated, were infected with Ad5 and Ad12 and levels of Tab182 monitored
227 (Figures 7C, 7D and 7E). In the Cul2-negative cells Tab182 is more stable following Ad12 infection
228 compared with the control cell line indicating that Cul2 is required for degradation of Tab182 by this
229 serotype (Figures 7D and 7C). In contrast, in the Cul5-negative cells Tab182 levels are comparable
230 with control cells following Ad12 infection indicating that this cullin component is dispensable for

231 Tab182 degradation (Figures 7E and 7C). However, more subtle differences were observed between
232 the degradation of Tab182, after Ad5 infection, in the Cul2-negative and Cul5-negative cells (Figures
233 7D and 7E). During Ad5 infection the loss of either cullin may result in some stabilization of Tab182
234 compared to control H1299 cells but does not clearly abrogate its degradation (Figures 7C, 7D and
235 7E). As expected MRE11 is stabilized in the Cul5-negative cells but not the Cul2-negative cells after
236 Ad5 infection. This suggests the possible involvement of Cul2, and/or perhaps an unidentified cullin,
237 in Ad5-mediated Tab182 degradation. Further work will be required to determine if other proteins
238 beside cullins 2 and 5 are involved in protein degradation by Ad5.

239 *Tab182 does not localize to viral replication centres*

240 It has previously been shown that a number of DDR proteins localize to the sites of adenovirus
241 replication in the nucleus, known as viral replication centres (VRCs) (6 and 8). To examine if this
242 applied to Tab182, HeLa cells were transfected with GFP-Tab182 and left for 24 hours. They were
243 then seeded onto glass coverslips and infected with Ad5 or Ad12. After a further 24 hours cells were
244 fixed and stained with antibodies that recognise VRCs (Figure 8). In the case of Ad5, VRCs were
245 visualised using an antibody against the viral DNA binding protein (DBP) while RPA-32 was used as a
246 surrogate marker for Ad12 VRCs. No specific recruitment of Tab182 to viral replication centres was
247 observed following infection with either adenovirus serotype (Figure 8A). As expression of GFP-
248 Tab182 was greater than is the case for the wt protein, in a further experiment soluble proteins
249 were extracted before antibody staining; again no co-localisation of GFP-Tab182 with VRCs was
250 observed (Figure 8B).

251 *Tab182 associates with AdE1B55K proteins.*

252 As the adenovirus-mediated degradation of Tab182 is AdE1B55K dependent we investigated
253 whether the two proteins associated, as is the case, for example, with Ad5E1B55K and p53 (57). To
254 examine this possibility, initially GST pull-down assays were carried out with purified GST-Tab182 (C-
255 terminal region) and whole cell lysates from E1B55K-expressing Ad12E1HER2 and Ad5E1HEK293
256 cells. In both cases the E1B55K protein was identified as a binding partner (Figures 9A and 9B). As

257 well as GST, GST-PRMT1 was included as an irrelevant (negative) control as it is of comparable
258 molecular weight to the Tab182 polypeptide. No binding of GST or GST-PRMT1 to E1B55K proteins
259 was seen. In further experiments, using the same cell lines, E1B55K proteins were co-
260 immunoprecipitated using an antibody against Tab182 (Figures 9C and 9D). No co-
261 immunoprecipitation was seen using an irrelevant antibody against collagen IV. In a further
262 experiment, Ad5E1HEK293 cells and Ad12E1HER2 cells were transfected with a construct encoding
263 GFP-Tab182. The lysates were immunoprecipitated with antibodies against AdE1B55K proteins and
264 co-precipitated Tab182 detected by western blotting (Figure 9E). The slightly higher molecular
265 weight GFP-Tab182 and the endogenous Tab182 were both seen in some lanes. These results
266 strongly suggest that, in both Ad5 and Ad12 serotypes, the viral E1B55K proteins interact directly
267 with Tab182.

268 Although degradation of Tab182 occurs to only a very limited extent during infection with
269 adenoviruses other than Ad5 and Ad12 (Figure 5) we considered the possibility that the E1B55K
270 proteins from these other species may also associate with Tab182. Therefore, HeLa cells were
271 transfected with constructs encoding HA-tagged Ad9E1B55K (group D) or HA-tagged Ad16E1B55K
272 (group B1). After 48 hours Tab182 was immunoprecipitated and associated E1B55K protein detected
273 with an antibody against HA (Figure 9F). It can be seen that whilst the Ad9 protein bound strongly
274 the Ad16 equivalent could only be seen on over-exposed western blots, indicating a very weak
275 association (Figure 9G). Similar results were obtained when the constructs were transfected into
276 Ad5E1HEK293 cells (data not shown). To check whether this differentiation extends to other
277 adenovirus targets the interaction with p53 was examined. After transfection of both constructs into
278 Ad5E1HEK293 cells, HA-tagged E1B55K proteins were immunoprecipitated and bound p53 detected
279 by western blotting (Figure 9H). In contrast to Tab182, both Ad9 and Ad16E1B55K proteins strongly
280 interacted with p53.

281 It is now well-established that the small DNA tumour viruses have many cellular targets in common,
282 such as pRb, p53 and CBP/p300 (58 and 59). It has already been reported that the E6 protein from

283 HPV genus beta, species 2 (HPV17a and HPV38) associates with the CNOT complex (60). To examine
284 if Tab182 interacts with proteins from other small DNA tumour viruses, a co-immunoprecipitation
285 experiment was carried out with Tab182 antibody using 293FT cells which express SV40 T antigen.
286 When Tab182 was immunoprecipitated appreciable SV40T was associated with it (Figure 9I).

287 *Tab182 is a component of the CNOT complex.*

288 It has previously been noted that Tab182 can be co-immunoprecipitated with the CNOT complex
289 from mammalian cells (40). To confirm this association, Tab182 was immunoprecipitated, using the
290 rabbit antibody raised against the C-terminal fragment, and the total immunoprecipitate analysed by
291 mass spectrometry. Results from a representative co-immunoprecipitation experiment are
292 presented in Table 1. In all cases most components of the CNOT complex were detected although
293 there were limited variations from one experiment to the next. Specifically, CNOT4 was never
294 detected in any Tab182 co-immunoprecipitate and CNOT 7 and CNOT 8 were occasionally not
295 identified. Significantly, neither Tab182 nor CNOT proteins were detected in any of the control
296 immunoprecipitates carried out with rabbit IgG (data not shown). Proteins which were seen in both
297 Tab182 and control IgG immunoprecipitations have not been listed in Table 1. The proteins listed are
298 the only ones which were consistently observed in five Tab182 immunoprecipitation experiments
299 but not in controls.

300 In a final set of co-immunoprecipitations we investigated whether AdE1B55K proteins were
301 associated with other CNOT components. Using the adenovirus E1-expressing cells, CNOT1 was
302 immunoprecipitated and associated E1B55K proteins detected by Western blotting (Figures 9J and
303 9K). It is possible that these results show direct interaction of the viral proteins with CNOT1 but
304 could also indicate interaction with other, as yet unidentified, components of the intact CNOT
305 complex or even Tab182.

306 *Adenovirus infection leads to a reduction in the levels of other CNOT proteins*

307 In light of the co-immunoprecipitation experiments shown in Table 1 and Figure 9 we examined the
308 levels of other CNOT proteins during adenovirus infection. Following infection of HeLa cells with

309 either Ad5 or Ad12 levels of CNOT1, CNOT3, CNOT4 and CNOT7 were monitored by Western blotting
310 (Figure 10) In contrast to Tab182, levels of CNOT1 and CNOT4 remained stable throughout the time
311 course of infection (Figures 10A and 10B). However, levels of both CNOT3 and CNOT7 were
312 markedly reduced after infection with both serotypes. In the case of Ad5, levels of CNOT3 decline
313 prior to the observed decrease in CNOT7 whereas for Ad12 CNOT7 levels decline prior to CNOT3
314 (Figures 10A and 10B). Further work will be required to determine whether these proteins are
315 degraded in the same fashion as Tab182 and whether levels of other CNOT proteins are reduced
316 during adenovirus infection but these data suggest that the complex may be a major target for
317 certain adenoviruses.

318 *Tab182 and CNOT1 depletion favours adenovirus infection*

319 To determine what advantage adenoviruses might derive from the degradation of Tab182 and other
320 CNOT complex proteins, a time course of infection was followed in HeLa cells treated with Tab182
321 siRNA. In addition, the effect of depletion of CNOT1 was also examined. CNOT1 forms a scaffold on
322 which other members of the complex associate (41). We, therefore, reasoned that its depletion
323 would cause maximal disruption of CNOT complex activity. Cells were infected with Ad5 and Ad12 48
324 hours after the addition of control, Tab182 or CNOT1 siRNAs. It can be seen from Figures 11A and
325 11B that, in the absence of Tab182, expression levels of the E1A viral proteins were elevated to a
326 limited extent compared to controls, during the time-course of infection. Similar results were
327 obtained with the E1B55K negative viruses, Ad5d/1520 and Ad12d/620, in that AdE1A proteins were
328 expressed at a higher level in the absence of Tab182 (data not shown). In the samples treated with
329 control siRNA there is a reduction in the level of Tab182 as degradation proceeds. Expression of
330 other viral proteins varied marginally between Tab182-depleted and control cells. However, in a
331 further set of experiments it was shown that when CNOT1 was depleted before infection with Ad5
332 there was a notable increase in E1A and E1B55K expression compared to controls (Figure 11A). In
333 Ad12 infected cells there was an even more marked increase in expression of the E1A and E1B55K
334 proteins, compared to control siRNA treated cells (Figure 11B). From the western blots it is clear

335 that Tab182 depletion has its most marked effect at 24 hours post Ad12 infection. However,
336 depletion of CNOT1 causes a several fold increase in Ad12E1A expression at 24 hours but, notably,
337 the level of protein stays consistently high up to 96 hours. The effects on Ad5E1A were less
338 pronounced, although, again, loss of Tab182 had most effect at 24 hours post infection whereas
339 depletion of CNOT1 facilitated AdE1A expression up to 96 hours.

340 It has long been known that adenovirus infection promotes cell cycle progression from G1 into a
341 'pseudo S-phase' accompanied by enhanced expression of cyclin E (reviewed in 61, for example). In
342 addition, it has also been reported that Ad E1A promotes expression of the tyrosine phosphatase
343 CDC25A, which is required for the G1 to S-phase transition (62). In an attempt to examine whether
344 the depletion of Tab182 and CNOT1 affects the ability of adenoviruses to initiate cell cycle
345 progression the expression of CDC25A was initially examined. It is notable that, in the Tab182 and
346 CNOT1 depleted cells, there is only very limited induction of CDC25A after infection whereas this is
347 appreciable in the control infected cells (Figure 11A and 11B). After 24 hours in all cases, expression
348 returns to a low level comparable with uninfected cells. We suggest that low level expression of
349 CDC25A is required by the virus for progression of the infected cells into pseudo S-phase but after
350 that, to stop further progression, CDC25A may be detrimental to viral replication. It is possible that
351 reduction in CNOT components decreases CDC25A, retaining the cells in a cell cycle phase more
352 conducive to viral early protein expression and viral replication.

353 *Tab182 depletion favours progression into S phase after adenovirus infection*

354 In view of the CDC25A western blotting data shown in Figure 11, the effects of depletion of Tab182
355 or CNOT1 on cyclin E expression were also examined. HeLa cells were again treated with control,
356 Tab182 and CNOT1 siRNAs, mock-infected or infected with Ad12 and then harvested at various
357 times up to 96 hours. In the mock-infected cells treated with control siRNA cyclin E is expressed at
358 constant low level but in those cells treated with Tab182 and particularly CNOT1 siRNAs there is an
359 appreciable elevation in cyclin E expression (Figure 12A). When a similar set of cells were infected
360 with Ad12 elevated cyclin E levels were also observed (Figure 12B). Thus, in control infected cells

361 there was a limited increase in cyclin E expression but in the absence of Tab182 or CNOT1,
362 expression of cyclin E is elevated to a much greater extent (Figure 12B). It seems likely, therefore,
363 that effects seen in the virally infected cells are primarily attributable to CNOT1 and Tab182
364 depletion rather than the virus itself. The advantage gained by the virus, facilitating E1A expression,
365 could be due to the fact that the siRNA treated cells have generally progressed slightly further
366 through the cell cycle, into a phase more favourable for adenovirus early protein expression, as
367 suggested in the previous section. Ad5 infection of HeLa cells treated with the same siRNAs had little
368 additional effect on cyclin E expression (data not shown).

369 *Tab182 and CNOT1 depletion enhances the AdE1A mRNA expression*

370 To determine whether depletion of Tab182 or CNOT1 affects AdE1A expression at the transcriptional
371 level, cells depleted of either Tab182 or CNOT1 were infected with either Ad5 or Ad12 before
372 isolation of total RNA after 24 hours. RT-PCR was performed following reverse transcription of total
373 RNA to amplify Ad5 and Ad12 13S E1As using primers across the CR3 unique region of each protein;
374 Ct values were calculated, normalised to GAPDH. Depletion of CNOT1 or Tab182 in Ad5 or Ad12
375 infected cells was verified by western blotting (data not shown). The relative expression of 13S E1A
376 in infected cells with depleted CNOT1 or Tab182 was compared with mock-transfected, infected
377 cells. It can be seen from the data presented in Figure 13 that depletion of Tab182 resulted in an
378 increase in both Ad5 and Ad12 13S E1A mRNAs compared to controls. The depletion of CNOT1 had a
379 more marked effect, consistent with the western blots shown in Figure 11.

380 *Tab182 and CNOT1 depletion favours the production of viral DNA during infection*

381 To examine whether the advantage gained in the expression of early proteins in Tab182 and CNOT1
382 depleted cells extends to the production of viral genomes HeLa cells were treated with appropriate
383 siRNAs and infected with Ad5 and Ad12 48 hours later. After a further 24 hours cells were harvested
384 and the DNA isolated. The concentration of adenovirus DNA was measured by quantitative PCR as
385 outlined in the Materials and Methods using primers equivalent to Hexon and GAPDH as a control.
386 More viral DNA can be seen in the Tab182 depleted cells than in the control cells after both Ad5 and

387 Ad12 infection (Figure 14); similarly, there is an even greater increase after CNOT1 depletion,
388 consistent with increased AdE1A expression. Interestingly, the effect of CNOT1 and Tab182
389 depletion were very similar during Ad5 infection (Figure 14A) whereas CNOT1 depletion had an
390 appreciably greater effect than Tab182 depletion in Ad12 infected cells (Figure 14B).

391

392 Discussion

393 It is now well-established that adenovirus infection triggers a cellular DDR (22). This is counteracted,
394 in Ad5 and Ad12 at least, by the degradation of multiple cellular proteins. Initially, it was noted that
395 p53 was a target for proteasome-mediated degradation during adenovirus infection (63 and 64). This
396 has been followed by demonstrations that other DDR proteins, such as MRE11, BLM and DNA Ligase
397 IV, are targeted to the proteasome through the actions of the viral E1B55K and E4orf6 proteins (10,
398 11 and 24). Whilst this is the case for the group A and group C viruses it certainly does not apply
399 universally to all adenovirus serotypes (28 and 56). In particular, it has been shown that group B (for
400 example, Ad7, Ad11 and Ad16) and group D (for example, Ad9) viruses target a much more limited
401 set of DDR proteins, possibly not extending beyond DNA Ligase IV. Furthermore, it seems that the
402 E1B55K/E4orf6 complex is not always required as degradation of TOPBP1 requires only Ad12E4orf6,
403 whereas DAXX degradation utilizes only Ad5E1B55K (12 and 27).

404 In a screen looking for additional DNA damage response proteins which might be targeted for
405 adenovirus-mediated degradation we have identified Tab182 and, subsequently other members of
406 the CNOT complex, CNOT7 and CNOT3, as probable targets. Tab182 was originally shown to interact
407 with tankyrase 1 (32) and to be highly phosphorylated by ATM and/or ATR after DNA damage by IR
408 (33). More recently, evidence has been presented to show that Tab182 plays a role in DSB repair and
409 promotes the association of PARP-1 with the DNA-PK catalytic subunit (34, 35).

410 There had been suggestions that Tab182 was a peripheral component of the CNOT complex in
411 mammals (40) and it was identified in various complexes in large scale protein interactome screens
412 (see, for example, 65-67). We have now confirmed that Tab182 is an integral component of the

413 CNOT complex. Depletion of the protein increases the sensitivity of cells to damage induced by
414 ionising radiation, UV radiation and HU and impairs the cell's ability to form DNA repair foci
415 following DNA replication stress (34, 35 and our unpublished data).

416 Here it has been shown that Tab182 is degraded during Ad5 and Ad12 infection (Figure 1). In both
417 cases this requires the AdE1B55K and AdE4orf6 proteins but is independent of AdE4orf3 which has
418 been shown to be required for degradation of other cellular proteins (25) (Figures 1 and 2).
419 Degradation of Tab182 is inhibited by bortezomib, a proteasome inhibitor, and MLN4924, which
420 inhibits cullin NEDDylation, preventing its activation (Figures 6 and 7). As is the case for p53
421 degradation, Ad12 hi-jacks a cullin 2-based E3 ligase (Figure 7), although it appears that ablation of
422 either Cul2 or Cul5 expression has a similar effect on Tab182 degradation during Ad5 infection
423 (Figure 7) in that loss of either causes partial protein stabilization. Clarification of this observation
424 requires further investigation.

425 To confirm the results of the mutant virus infections, that Tab182 is targeted through AdE1B55K, co-
426 immunoprecipitation assays were carried out and it was found that Tab182 and both the Ad5 and
427 Ad12 proteins could be immunoprecipitated together. Furthermore, both Ad5 and Ad12 E1B55K
428 proteins bound to the GST-Tab182 C-terminal region, indicating a direct interaction (Figure 9).
429 Interestingly, Tab182 binds strongly to Ad9E1B55K but not Ad16E1B55K although it does not appear
430 to be degraded by either group B1 (Ad7 and Ad16) or group D (Ad9) viruses (Figures 5 and 9).
431 E1B55K proteins from both Ad9 and Ad16 interact with p53 as might be expected since it is
432 transcriptionally inactive after Ad9 and Ad7 infection, even though it is present at high level (28).
433 These observations suggest that interaction of E1B55K with Tab182 may be determined by factors
434 other than a requirement for protein degradation. A more widespread examination of the
435 interaction of Tab182 with E1B55K proteins from a number of adenoviruses may elucidate this point.
436 Tab182 also associates with SV40T antigen in co-immunoprecipitation experiments, suggesting that
437 the protein could be a target for the family of small DNA tumour viruses (Figure 9I). Significantly,

438 previous studies have shown that the CNOT complex associates with HPV17a and HPV38 E6 proteins
439 although the consequences of this for the virus were not examined at the time (60).
440 To see how extensive the relationship between adenoviruses and the CNOT complex was, the fate of
441 other components of the complex was studied following adenovirus infection (Figure 10). Although
442 only a limited number of CNOT proteins were examined it was seen that levels of CNOT3 and CNOT7
443 were reduced during Ad5 and Ad12 infection whereas levels of CNOT4 and CNOT1 remained stable.
444 Although a number of activities have been attributed to the CNOT complex, such as deadenylase,
445 transcriptional regulation and ubiquitin E3 ligase activity (37-39 and 46-48) it is not clear what
446 contribution Tab182 makes. To attempt to understand why adenovirus might target Tab182 (and
447 other CNOT proteins) adenovirus infection was compared in control and siRNA knock down cells. It
448 was seen that expression of E1A was enhanced, to a limited extent, in Tab182 depleted cells
449 although little or no difference was seen in the expression of late proteins (Figure 11). To see if a
450 similar effect occurred with other members of the CNOT complex, CNOT1, which is considered to be
451 a scaffold protein required for the integrity of the complex, was depleted. During Ad5 infection E1A
452 expression was notably increased when CNOT1 was depleted while in the case of Ad12 there was a
453 greatly enhanced expression of E1A and a marked increase in E1B55K protein, following CNOT1
454 knock down, compared to controls (Figure 11). The increased effect of CNOT1 protein depletion on
455 Ad12 compared to Ad5 appears to be consistent with Ad12's somewhat enhanced ability to degrade
456 Tab182. The difference in expression of E1A protein is due to an increase in AdE1A mRNA, as shown
457 by RT-PCR (Figure 13). Whether this effect is directly attributable to a reduction in deadenylase
458 activity of the CNOT complex will have to await further investigation. Interestingly, it has recently
459 been shown that the Ad5 E1B55K/E4orf6 complex enhances E1A activity by stabilizing the protein,
460 leading to increased level, and by increasing the activation of E2F by E1A (68). It is possible that the
461 effect of the same adenovirus complex on the CNOT complex, as demonstrated here, could
462 contribute to the increased AdE1A level observed. In a further study it has been shown that the
463 concentration of viral DNA is increased in Tab182 and CNOT1 depleted cells 24 hours after both Ad5

464 and Ad12 infection (Figure 14). More marked effects were seen with CNOT1 depletion than with
465 Tab182 consistent with the observed increase in AdE1A expression (Figure 11); however, reduction
466 in Tab182 had less effect on relative Hexon DNA concentration after Ad12 infection than with Ad5,
467 reasons for this are not clear at present.

468 The relationship between adenoviruses and the CNOT complex is not clear cut, for whilst the virus is
469 able to cause degradation of various components this occurs later than any initial enhanced increase
470 in AdE1A expression seen after the depletion of CNOT proteins described here. It is notable that
471 there is a sustained increase in AdE1A expression up to 96 hours in the absence of CNOT1 (Figure
472 11). However, increases in viral DNA concentration were observed after CNOT1 and Tab182
473 depletion, suggesting that inactivation of the complex will facilitate viral replication to a limited
474 extent. It is also possible that the aim of the virus, in degrading and presumably inactivating the
475 CNOT complex, is not necessarily just to facilitate AdE1A expression, but to fulfil some other, as yet
476 unidentified, role, perhaps linked to an effect on the DDR. It should be borne in mind, when
477 considering the effects of CNOT1 depletion, that adenoviruses do not actually cause its degradation
478 and while its loss will probably indicate the effect of inactivation of the CNOT complex it does not
479 necessarily coincide with what happens *in vivo*. It is also possible that other CNOT proteins could be
480 targets for adenovirus-mediated degradation early in infection co-incident with AdE1A expression,
481 although we have no evidence of this. Significantly, degradation of Tab182 and CNOT7 occurs later in
482 viral infection than is the case for MRE11 and BLM and is more similar to that seen for p53.

483 Loss of components of the CNOT complex, for example Tab182, appears to facilitate progression of
484 cells into late G1/early S-phase, as evidenced by the enhanced expression of cyclin E and transiently
485 enhanced expression of CDC25A (Figures 11 and 12). This may provide an environment more
486 conducive to expression of early viral proteins, particularly E1A. For reasons which are not evident at
487 present the effect seems to be more marked with Ad12 compared to Ad5. With relevance to the
488 effects on cyclin E expression, it is worth noting that CNOT1 depletion has a more marked effect

489 than does Tab182 depletion, suggesting that its loss enhances cell cycle progression to a greater
490 extent. However, it is possible that loss of other CNOT proteins could have a comparable effect.

491 In summary, Ad5 and, in particular, Ad12 have been shown to target Tab182 and other CNOT
492 proteins for proteasome-mediated degradation during viral infection. Loss of Tab182 and CNOT1
493 favours enhanced expression of AdE1A and E1B55K proteins in the early stages of infection.

494 **Materials and methods**

495 *Cell lines, viruses and plasmids.*

496 HeLa (obtained from ATCC), HEK293FT (Invitrogen), Ad5E1HEK293 (a generous gift from Frank
497 Graham), and Ad12E1HER2 (69) cells were grown in DMEM supplemented with 8% foetal calf serum
498 (FCS). H1299-based cell lines in which Cul2 or Cul5 expression had been ablated were a generous gift
499 from Paola Blanchette and Phil Branton. The cells were grown in DMEM supplemented with 8% FCS
500 and 1 µg/ml puromycin (Cul2⁻) or 8 % FCS, 1 µg/ml puromycin and 100 µg/ml hygromycin (Cul5⁻).
501 Ad4, Ad5, Ad7, Ad9, Ad11 and Ad12 viruses were obtained from ATCC or were a generous gift from
502 Jo Mymryk. The following Ad5 mutant viruses were used: Ad5dl1520 (Ad5E1B55K⁻) (70), H5in351
503 (E4orf1⁻), H5pm4154 (E4orf6⁻), H5pm4155 (E4orf3⁻, E4orf6⁻), H5pm4166 (E4orf4⁻), H5dl356 (E4orf7⁻)
504 and H5in352 (E4orf2⁻) (23, 26, 71-73). In addition, an Ad12 E1B55K negative mutant virus
505 (Ad12dl620) was used (74). HeLa cells were generally infected at a multiplicity of infection of 5
506 plaque forming units (pfu)/cell. Ad5 and Ad12 E1B55K DNA was cloned into pcDNA3 and Ad5 and
507 Ad12 E4orf6-HA tag DNA was also cloned into pcDNA3 as previous described (75). NEDDylation was
508 inhibited by addition of MLN4924 to the cell culture medium at a concentration of 4 µM and
509 proteasomal activity was inhibited with bortezomib (0.5 µM).

510 *siRNA treatment to deplete Tab182 and CNOT proteins and protein transfections.*

511 HeLa cells were plated at a density of 4x10⁵ per 6 cm dish. After 24 hours they were transfected with
512 control or ON-TARGETplus SMART pool siRNAs (0.2 nmol/dish) (GE Dharmacon) directed against
513 Tab182 or CNOT1 proteins using Oligofectamine (Invitrogen) following the manufacturer's protocol.
514 After 24hours cells were split 1→3 and after a further 24hours infected with virus. For protein

transfections cells were grown to 70% confluency and then incubated with DNA constructs (2µg/6cm dish or 5µg/10cm dish) which had been previously mixed for 20 minutes with Lipofectamine 2000 (Invitrogen) in Opti-Mem (Gibco) following the manufacturer's protocol. After 24 hours cells were incubated with fresh medium and harvested 24 hours later.

Cloning *Tab182*

Total cellular RNA was isolated from a lymphoblastoid cell line from a normal individual using the Qiagen RNeasy Mini Kit and was reverse transcribed into cDNA using the oligo-dT primer d(T)23VN and the Protoscript II First Strand cDNA Synthesis Kit (New England Biolabs). PCR was used to amplify the complete *Tab182* cDNA sequence using the forward primer (For 1) 5'-GAGCGGGTCGACGATGAAAGTGTCTACTCTCAGG-3' and the reverse primer (Rev13) 5'-CGTGATGTCGACTCAGACCTTCTTCTTCTTCAGTTT-3'. Both primers contain the recognition sequence for the restriction enzyme *Sal* I (underlined). The forward primer contains the translation initiation codon for *Tab182* (italics) and the reverse primer contains the translation termination codon for *Tab182* (italics; strand antiparallel to sense strand). The *Tab182* cDNA sequence was amplified using Q5 High-Fidelity DNA Polymerase (New England Biolabs). An initial denaturation step of 98°C for 30 seconds was followed by 30 cycles of 98°C for 5 seconds, 62°C for 15 seconds and 72°C for 4 minutes. A final extension of 5 minutes at 72°C followed the 30 cycles. The PCR products were analysed by gel electrophoresis and a product of the correct size (5190 base pairs) was identified. The products were digested with *Sal* I-HF and the excised *Tab182* band purified by gel electrophoresis. *Tab182* was cloned into the pEGFP-C3 plasmid. Sequence determination was performed using an Applied Biosystems 3500xL Genetic Analyzer. Sequences were analysed on-line using BLAST at the National Center for Biotechnology Information (NCBI). Sequences were all wild type. Codon 322 can encode threonine (ACT) or serine (AGT) and the ratio is approximately equal in the general population. The sequences isolated from the individual used to make the cDNA for this cloning exercise were all found to encode serine at amino acid 322.

Isolation of RNA and cDNA Synthesis

541 Cellular RNA was extracted using the SV Total RNA Isolation System (Promega) following the
542 manufacturer's protocol. To remove any DNA contamination, RNA was treated with DNase I
543 (Promega). RNA quantity and quality were evaluated by optical density measurements (260/280 nm
544 ratios) and by agarose gel electrophoresis. First-strand cDNA synthesis was performed using
545 SuperScript™ II Reverse Transcriptase (RT) (Invitrogen) and random primers according to the
546 manufacturer's instructions.

547 *Isolation of genomic DNA*

548 Cellular DNA was extracted using the QIAamp DNA Mini Kit (Qiagen) following the manufacturer's
549 protocol. In order to remove any protein or RNA contamination 15 µl Proteinase K (10 mg/ ml)
550 (Sigma-Aldrich) and 4 µl RNase A (20 mg/ml) (Invitrogen) were added to each sample. DNA quantity
551 and quality were evaluated by optical density measurements (260/280 nm ratios) and by agarose gel
552 electrophoresis.

553 *Primer design and RT-PCR*

554 Cellular RNA or DNA was extracted as described above. The sequences of the primers used for RT-
555 PCR are as shown in Table 2. Specificity of the primers was checked with NCBI/Primer-BLAST.
556 The RT-PCR reactions were performed in the Mx3005P system (Stratagene) using real-time
557 PowerUp™ SYBR® Green Master Mix (Applied Biosystems). Quantitative RT-PCR was carried out in a
558 final volume of 20 µl containing 2 µg or 10ng of cDNA or DNA, respectively, 5 pmol of the forward
559 primer, 5 pmol reverse-primer and 10 µl of PowerUp™ SYBR® Green Master Mix. Thermocycling
560 program was performed for 10 min at 95°C for the pre-cycling step to denature the cDNA and to
561 activate Dual-Lock™ Taq DNA Polymerase, and then followed by 35 cycles of denaturation at 95°C
562 for 30 sec, annealing at 55°C for 1 min and extension at 72°C for 1 min. To confirm the expected
563 amplifications 2% agarose gel electrophoresis with ethidium bromide staining was performed. Viral
564 AdE1A or Hexon and host cell Tab182 and CNOT1 Ct values were normalized to Ct values of GAPDH

565 amplified from the same sample [for example, $\Delta Ct = Ct \text{ (Tab182)} - Ct \text{ (GAPDH)}$], and the $2^{-\Delta\Delta Ct}$
566 method was used to calculate the relative-expression. Each experiment was performed in triplicate.

567 *Western blotting and antibodies.*

568 Cells were harvested after washing with ice-cold phosphate buffered saline (PBS) and solubilised in 8
569 M urea, 50 mM Tris HCl pH7.4, and 0.15 M β -mercaptoethanol. Proteins were fractionated on
570 polyacrylamide gels in the presence of 0.1 M Tris, 0.1 M Bicine, and 0.1% SDS. For western blotting,
571 proteins were electrophoretically transferred to nitrocellulose membranes before incubation with
572 antibodies overnight at 4°C. Antibodies used in the study were as follows: Tab182 (an antibody
573 raised in rabbits against GST-Tab182 [C-terminal fragment]), MRE11, CNOT3, CNOT4, CNOT7, (all
574 from GeneTex), CNOT1 (Proteintech), cullin2, cyclin E1, RPA32 (Abcam), p53 (raised in rabbits),
575 cullin5, GAPDH, collagen IV, SV40T (Santa Cruz Biotechnology), and β actin (Sigma-Aldrich). Rabbit
576 antibodies against Ad5 Hexon and Ad12 Fiber protein were gifts from Vivien Mautner and Paul
577 Freimuth, respectively. A mouse monoclonal antibody against Ad5DNA binding protein (DBP) was a
578 gift from Pieter van der Vliet. Antibodies against Ad5E1A (M73), Ad12E1A (5DO2), Ad12E1B55K
579 (XPH9), Ad5E1B55K (2A6), p53 (DO1) and HA (12CA5) were purified from monoclonal supernatants.

580 *GST pull-down assays and co-immunoprecipitation*

581 The C-terminal fragment of Tab182 (amino acids 824-867+1221-1729) was expressed in *E.coli* as a
582 GST fusion protein as described (49). For GST pull-down and co-immunoprecipitation assays cells
583 were harvested in ice-cold PBS and lysed in 0.4 M NaCl, 40 mM Tris HCl pH7.4, 5 mM EDTA, 1%
584 NP40. Insoluble protein was removed by centrifugation (45K, 30 minutes, 4°C). Lysates were
585 incubated overnight either with GST fusion protein or appropriate antibody. Protein complexes were
586 retrieved on glutathione-agarose beads or Protein G-agarose beads as appropriate. After washing
587 with lysis buffer, bound proteins were released with either 25mM glutathione, pH8.2 (GST fusion
588 proteins) or SDS sample buffer (immunoprecipitated samples) and fractionated by SDS-PAGE prior to
589 western blotting.

590 *Mass spectrometry*

591 Proteins were immunoprecipitated as described above except that the antibody-antigen complexes
592 were released with 8 M urea, 50 mM NH_4HCO_3 for 30 minutes at ambient temperature. Proteins
593 were reduced in 50 mM DTT, 50 mM NH_4HCO_3 at 56°C for 30 minutes and then carboxymethylated
594 in 100 mM iodoacetamide at ambient temperature in the dark for 30 minutes. Proteins were
595 retrieved using Amicon centrifugal filters (30K molecular weight cut off) which were washed four
596 times with 50 mM NH_4HCO_3 . The filters, with the bound immunoprecipitated proteins, were
597 incubated overnight at 37°C with trypsin (1 μg) in 50 mM NH_4HCO_3 . Tryptic peptides were retrieved
598 by centrifugation, dried and analysed using a Bruker amaZon ion trap mass spectrometer. Peptides
599 were identified using the ProteinScape central bioinformatics platform (Bruker).

600 *Immunofluorescence microscopy*

601 HeLa cells were grown on glass cover slips. After 24 hours cells were infected or mock infected with
602 Ad5 or Ad12 (5pfu/cell) for 30 hours. Cells were fixed in 3.6% para-formaldehyde in PBS for 10
603 minutes and permeabilized in 0.5% TritonX-100 in PBS for 5 minutes. Fixed cells were stained with
604 primary antibodies for 1 hour, washed three times in PBS and stained with secondary antibodies also
605 for 1 hour. DNA was stained with DAPI. When pre-extraction was used cells were treated with pre-
606 extraction buffer (10 mM PIPES, 20 mM NaCl, 3 mM MgCl_2 , 300 mM sucrose, 0.5% Triton X-100) for
607 7 minutes on ice before fixing with 3.6% para-formaldehyde and antibody staining as above.
608 Fluorescence images were taken using a Nikon E600 Eclipse microscope³³³ equipped with a 60X oil
609 lens, and images were acquired and analysed using Volocity Software 334 v4.1 (Improvision).

610

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799

800 **Figure legends**

801 **Figure 1: The Degradation of Tab182 following infection with adenovirus serotype 5 or adenovirus**
802 **serotype 12 is dependent on the adenovirus E1B55K protein.** HeLa cells were infected with
803 adenovirus serotype 5 (A), or serotype 12 (B) at 5 pfu/cell. HeLa cells were also infected with
804 adenovirus serotype 5 E1B55K negative virus Ad5dl1520 (C), and adenovirus serotype 12 E1B55K
805 negative virus Ad12dl620 (D) at 10 pfu/cell. Cells were then harvested at various time points (0, 8,
806 24, 48, 72 and 96 hours) post-infection. Cell lysates were subjected to SDS-PAGE and Western
807 blotting using the indicated antibodies.

808 **Figure 2: The Degradation of Tab182 following infection with adenovirus serotype 5 is dependent**
809 **on the adenovirus E4orf6 protein.** HeLa cells were infected with Ad5 E4 mutants H5in351 (E4orf1-)
810 (A), H5pm4154 (E4orf6-) (A), H5pm4155 (E4orf3-E4orf6-) (B), H5pm4166 (E4orf4-) (B), H5dl356
811 (E4orf6-E4orf7-) (C) H5in352 (E4orf2-) (C) and H5pm4150 (E4orf3-) (D) at 10 pfu/cell. Cells were then
812 harvested at various time points (0, 8, 24, 48, 72 and 96 hours) post-infection. Cell lysates were
813 subjected to SDS-PAGE and Western blotting using the indicated antibodies.

814 **Figure 3: Tab182 gene expression is enhanced in adenovirus infected cells.** HeLa cells
815 were infected with Ad5 or Ad12 at 5 pfu/cell. Cells were harvested at various time points (0,
816 8, 24, 48, 72 and 96 hours) post-infection. Cellular RNA was extracted from Ad5 (A) and
817 Ad12 (B) infected cells and first-strand cDNA synthesis carried out. The RT-PCR reactions
818 were performed using Tab182-specific primers and real-time PowerUp™ SYBR® Green
819 Master Mix. To determine the relative Tab182 gene expression, calculated Tab182 Ct values
820 were normalized to Ct values of GAPDH amplified from the same sample [$\Delta Ct = Ct (\text{Tab182})$]

821 – Ct (GAPDH)], and the $2^{-\Delta\Delta Ct}$ method was used to calculate relative expression. Each
822 experiment was performed in triplicate. . Western blots of the Ad5 and Ad12 infected HeLa cells
823 were performed to confirm Tab182 degradation (data not shown).
824

825 **Figure 4: The Degradation of Tab182 during adenovirus serotype 5 and 12 Infection is dependent**
826 **on the adenovirus E1B55K and E4orf6 proteins.** 2 μ g of plasmid DNA as shown was transfected into
827 HeLa cells and 48 hours later cells were harvested and subjected to SDS-PAGE and Western blotting
828 using the indicated antibodies. Ad5 and Ad12 E4orf6 proteins were detected with an antibody which
829 recognised the HA tag. GAPDH is included as a loading control.

830 **Figure 5: Tab182 level following infection by Group B, D and E adenoviruses.** HeLa cells were
831 infected with: (A) Ad5 (group C) and Ad12 (group A), (B) Ad4 (group E) and Ad9 (Group D), and (C)
832 Ad11 (group B2) and Ad7 (group B1) at 5 pfu/cell. Cells were harvested at 8, 24, 48, 72, 96 and 120
833 hours post infection. Cell lysates were subjected to SDS-PAGE and Western blotting using antibodies
834 against Tab182, MRE11, p53 and β -actin. Hexon expression was confirmed, as a marker of viral
835 infection, by Ponceau S staining of Western blots for total protein.

836 **Figure 6: The down-regulation of Tab182 protein levels during Ad5 and Ad12 infection can be**
837 **rescued by the proteasomal inhibitor Bortezomib.** HeLa cells were infected with Ad5 (A) or Ad12 (B)
838 at 5 pfu/cell. Cells were treated with 0.5 μ M Bortezomib or DMSO control and harvested after 48
839 hours. Cell lysates were subjected to SDS-PAGE and Western blotting using the indicated antibodies.

840 **Figure 7: The Degradation of Tab182 during Ad5 and Ad12 infection is dependent on cullin**
841 **function.** HeLa cells were infected with Ad5 and Ad12 at 5 pfu/cell. Cells were treated with the
842 Nedd8 inhibitor MLN4924 (4 μ M) 1 hour before infection and retreated immediately post-infection.
843 Cells were harvested at various time points (0, 8, 24, 48, 72 and 96 hours) post-infection. Cell lysates
844 were subjected to SDS-PAGE and Western blotting using the indicated antibodies (A) and (B). H1299
845 cells (C) or H1299 cells with ablation of Cul2 (D) or Cul5 (E) expression were infected with either Ad5

846 or Ad12 and harvested at 0, 8, 24, 48, 72 and 96 hours post-infection. Cell lysates were subjected to
847 SDS-PAGE and Western blotting with the antibodies shown.

848 **Figure 8: Tab182 does not localise to viral replication centres during adenovirus infection.** GFP-
849 Tab182 was transfected into HeLa cells and 24 hours later cells were infected with Ad5 or Ad12. (A),
850 30 hours later cells were fixed, extracted and probed with the appropriate antibodies. (B) 30 hours
851 after infection cells were pre-extracted as described in the Materials and Methods section before
852 fixing and then staining with antibodies. In both (A) and (B) Ad5 infected cells were probed with DBP
853 antibody, whilst Ad12 infected cells were probed with RPA32 antibody. Nuclear DNA is stained with
854 DAPI.

855 **Figure 9: Adenovirus early region E1B55K interacts with Tab182 *in vitro* and *in vivo*.** Ad12E1HER2
856 (A) and Ad5E1HEK293 (B) cell lysates containing 500 µg total protein were incubated with 5 µg
857 either GST-Tab182, GST-PRMT1 or with GST alone. Protein complexes were captured by glutathione-
858 agarose beads, subjected to SDS-PAGE and Western blotting with the antibodies indicated.
859 Ad5E1HEK293 (C) and Ad12E1HER2 (D) cell lysates (500 µg total protein) were incubated with
860 antibodies against Tab182, collagen IV together with IgG (non-specific binding controls). Immuno-
861 complexes were isolated using Protein-G agarose beads and subsequently resolved by SDS-PAGE and
862 Western blotting using antibodies against Ad5/Ad12 E1B55K proteins. (E) GFP-Tab182 was
863 transfected into Ad5E1HEK293 and Ad12E1HER2 cell lines which were harvested after 48 hours. Cell
864 lysates (500µg total protein) were incubated with Ad5 and Ad12 E1B55K antibodies together with
865 IgG. Western blotting was with an antibody against Tab182. (F) HeLa cells were transfected with
866 pcDNA3 or pcDNA3 constructs expressing HA-tagged Ad9E1B55K or Ad16E1B55K. After 48hours
867 lysates (500µg total protein) were immunoprecipitated with an antibody against Tab182 or rabbit
868 IgG. Western blotting was with an antibody against HA. (G) is an over-exposed version of a portion of
869 the western blot shown in (F). (H) Ad5E1HEK293 cells were transfected with pcDNA3 or pcDNA3
870 constructs expressing HA-tagged Ad9E1B55K or Ad16E1B55K. After 48hours lysates (500µg total
871 protein) were immunoprecipitated with an antibody against HA or mouse IgG. Western blotting was

872 with an antibody against p53. (I) 293FT cell lysates (500µg protein) were incubated with antibodies
873 against Tab182, collagen IV or IgG control. Western blotting was with an antibody against SV40 T
874 antigen. Ad5E1HEK293 (J) and Ad12E1HER2 (K) cell lysates (500 µg total protein) were incubated
875 with antibodies against CNOT1, collagen IV together with IgG. Western blotting was with antibodies
876 against Ad5/Ad12 E1B55K proteins. In all cases the whole cell lysates contained 15 µg of protein.
877 Although only limited areas of the western blots are shown no additional bands were seen in the
878 original autoradiographs.

879 **Figure 10: Adenoviruses 5 and 12 degrade components of the CNOT complex.** HeLa cells were
880 infected with adenovirus serotype 5 (A) or 12 (B) at 5 pfu/cell. Cells were harvested at 0, 8, 24, 48,
881 72 and 96 hours post infection, subjected to SDS-PAGE and Western blotting using the indicated
882 antibodies.

883 **Figure 11: AdE1A protein expression is enhanced in adenovirus-infected, Tab182- or CNOT1-**
884 **depleted cells.** HeLa cells were transfected with control, Tab182 or CNOT1 siRNAs. 48 hours later,
885 control, Tab182 and CNOT1 siRNA treated cells were infected with adenovirus serotype 5 (A) or
886 serotype 12 (B) at 5 pfu/cell. Cells were then harvested at various time points (0, 8, 24, 48, 72 and 96
887 hours) post-infection. Cell lysates were subjected to SDS-PAGE and Western blotting using the
888 indicated antibodies.

889 **Figure 12: Expression of cyclin E and is enhanced in Tab182 and CNOT1 depleted cells.** HeLa cells
890 were transfected with control, Tab182 or CNOT1 siRNAs. 48 hours later, control, Tab182 and CNOT1
891 siRNA treated cells were mock infected (A) or infected with adenovirus serotype 12 (B) at 5 pfu/cell.
892 Cells were then harvested at various time points (0, 8, 24, 48, 72 and 96 hours) post-infection. Cell
893 lysates were subjected to SDS-PAGE and Western blotting using the indicated antibodies.

894 **Figure 13: The relative expression of Ad13S E1A mRNA is increased in infected cells in the absence**
895 **of CNOT1 or Tab182.** HeLa cells were transfected with control, Tab182 or CNOT1 siRNAs and 48
896 hours later infected with Ad5 (A) or Ad12 (B) at 5 pfu/cell. Cellular RNA was extracted from infected
897 cells and first-strand cDNA synthesis carried out. The RT-PCR reactions were performed using

Ad13SE1A CR3 region specific primers and real-time PowerUp™ SYBR® Green Master Mix. To check E1A relative gene expression, calculated E1A Ct values were normalized to Ct values of GAPDH amplified from the same sample [$\Delta Ct = Ct (E1A) - Ct (GAPDH)$], and the $2^{-\Delta\Delta Ct}$ method was used to calculate relative gene expression. Data are the mean of 3 repeats. The statistical significance was determined using Student's *t*-test, p-values less than 0.05 (*) or 0.01 (**) were considered significant. Error bars represent SEM.

Figure 14: Viral DNA synthesis is increased in Tab182 and CNOT1-depleted cells after adenovirus infection. HeLa cells were treated with control, Tab182 and CNOT1 siRNA for 48 hours and then infected with Ad5 (A) or Ad12 (B) at 5 pfu/cell. After 24 hours cells were harvested and the total DNA isolated. Quantitative PCR was performed to determine relative concentration of viral DNA. Hexon Ct values were normalized to Ct values for GAPDH DNA amplified from the same sample. Data are the mean of 3 repeats. The statistical significance was determined using Student's *t*-test, p-values less than 0.05 (*) or 0.01 (**) were considered significant. Error bars represent SEM.

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916 **Legend to Table 1**

917 **Table1 Proteins identified by mass spectrometric analysis after co-immunoprecipitation with**
918 **Tab182 antibody.** HeLa cells were immunoprecipitated with a rabbit antibody raised against the C-terminal fragment of Tab182 and analysed as described in the Methods section. These data are
919 representative of five independent experiments.
920

921 **Table 2 Primers used in this study**

922

923 **Table1**

Protein	Peptide	Percentage	Mascot Score
---------	---------	------------	--------------

	number	coverage	
Tab182	68	49.4	3491
CCR4-NOT1	38	17.4	1472
CCR4-NOT3	7	10.2	237
CCR4-NOT7	6	28.8	211
CCR4-NOT2	5	12.2	245
CCR4-NOT6L	1	1.8	21
CCR4-NOT10	1	1.3	29
C2orf29 (NOT11)	1	2.4	56
RCD1 (NOT9)	6	20.4	224
PRMT3	9	18.5	377
FHL2	12	52.3	470

924

925 **Table 2**

Gene	Sequence (5'→3') Forward primer Reverse primer	Length	Start	Stop	Product Size (bases)
E1A-Ad5 (CR3)	TAGATTATGTGGAGCACCCCG GCCACAGGTCTCATATAGCAA	21	990	1010	110
		22	1099	1078	
E1A-Ad12 (CR3)	AGTCCTGTGAGCACCACCG GTAGGCTCGCAGATAGCACA	19	980	1053	74
		20	998	1034	
Tab182	CTGCTCTGAGGGACTCCTTG CTGGGTCTCCTCTAGGGCTT	20	2310	2329	158
		20	2448	2467	
GAPDH (RNA)	GAGTCAACGGATTGGTCGT ACAAGCTTCCGTTCTCAG	20	53	72	183
		19	218	236	
GAPDH (DNA)	CGGCTACTAGCGGTTTTACG AGAAGATGCGGCTGACTGT	20	6534369	6534388	188
		20	6534538	6534557	
Hexon-Ad5	GCCACGGTGGGGTTTCTAACTT GCCCCAGTGGTCTTACATGCACATC	23	18862	18882	127
		25	18967	18989	
Hexon-Ad12	GCCACGGTGGGGTTTCTAACTT GCCCCAGTGGTCTTACATGCACATC	23	17764	17784	127
		25	17869	17891	

926

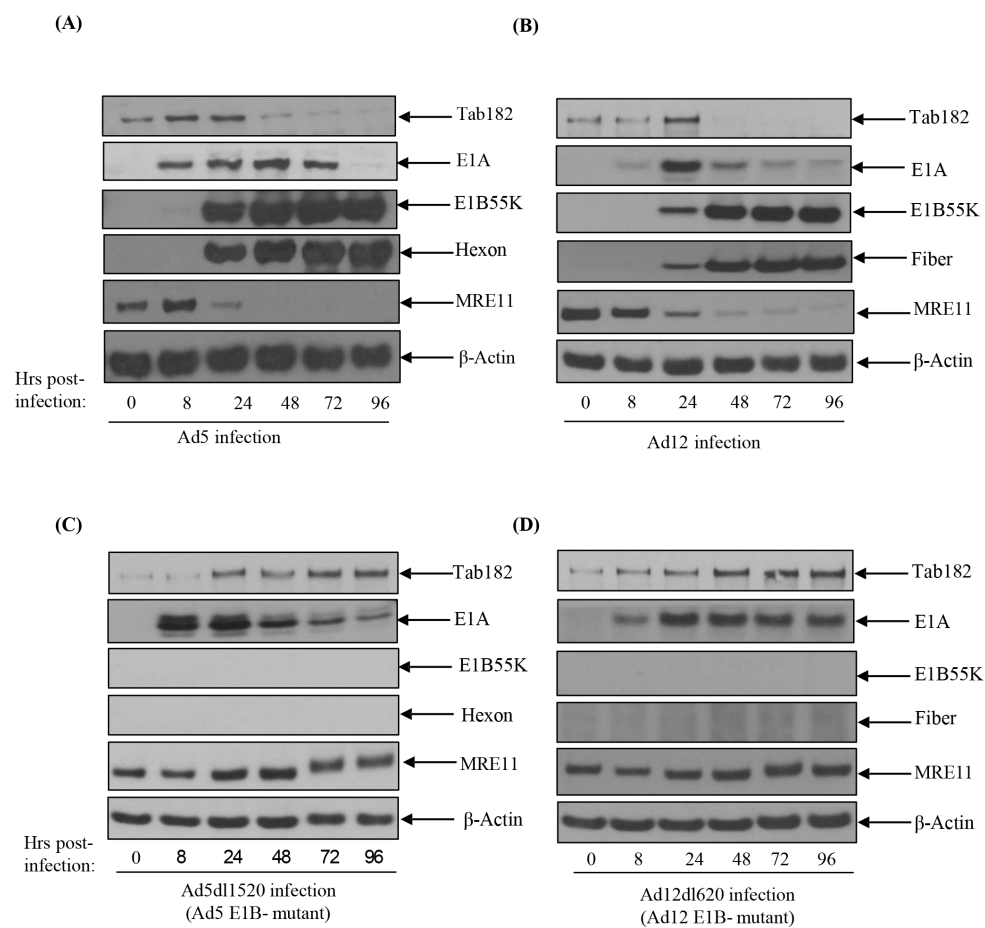


Figure 1

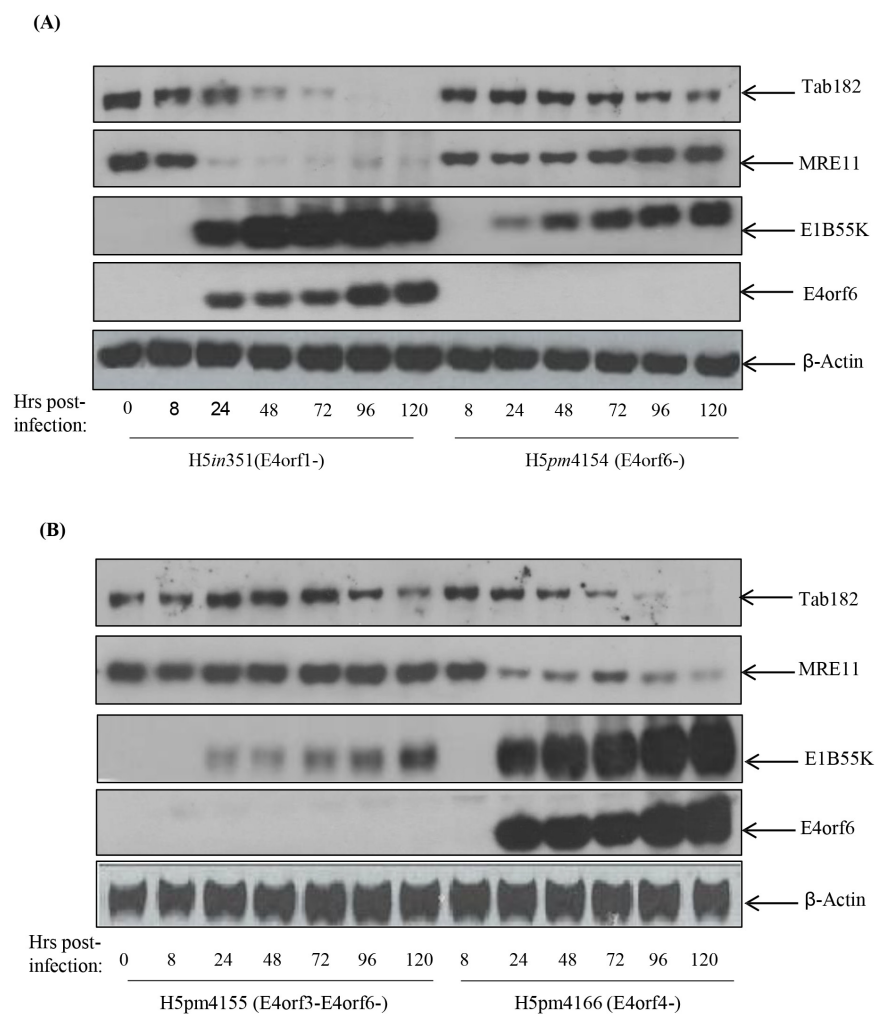


Figure 2

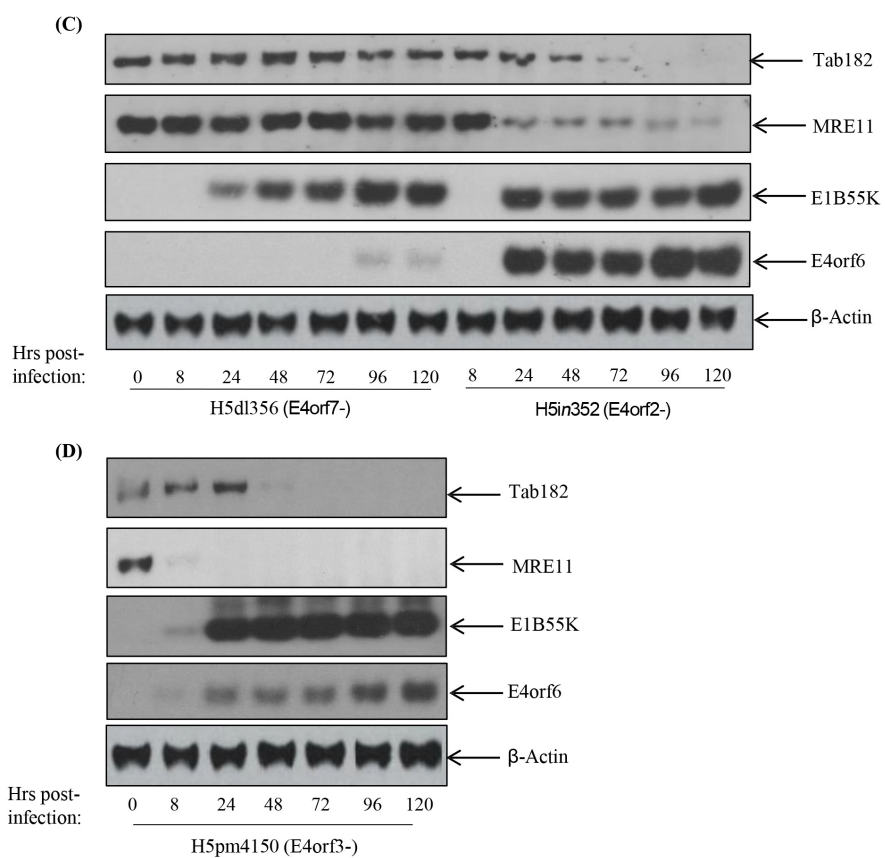


Figure 2

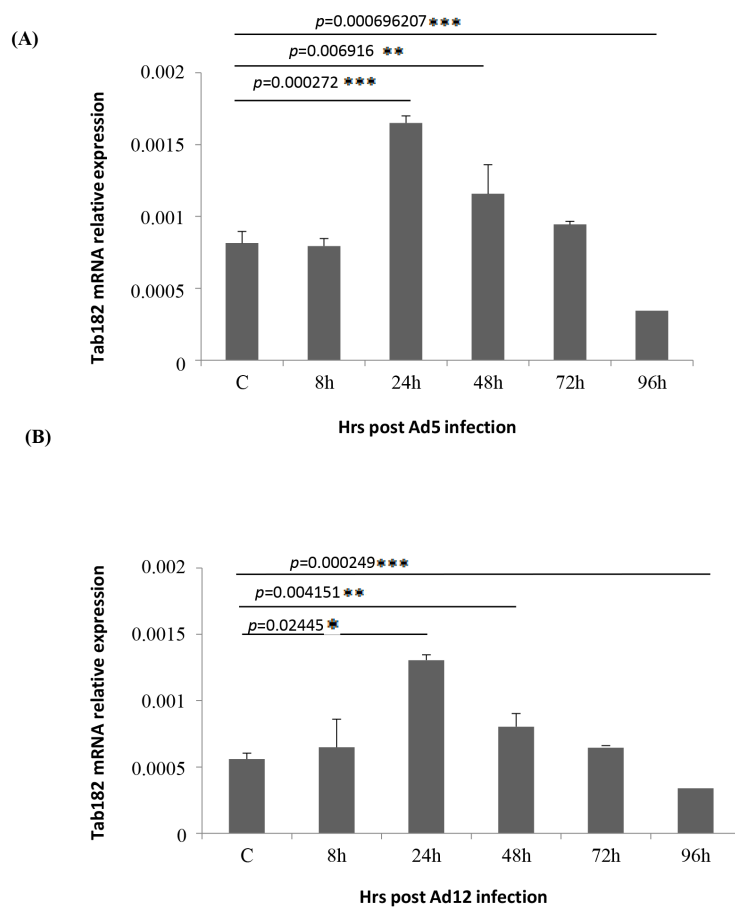


Figure 3

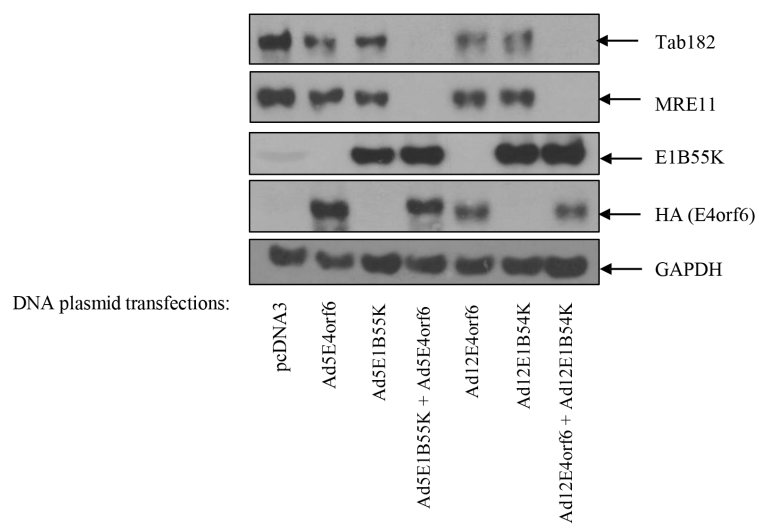


Figure 4

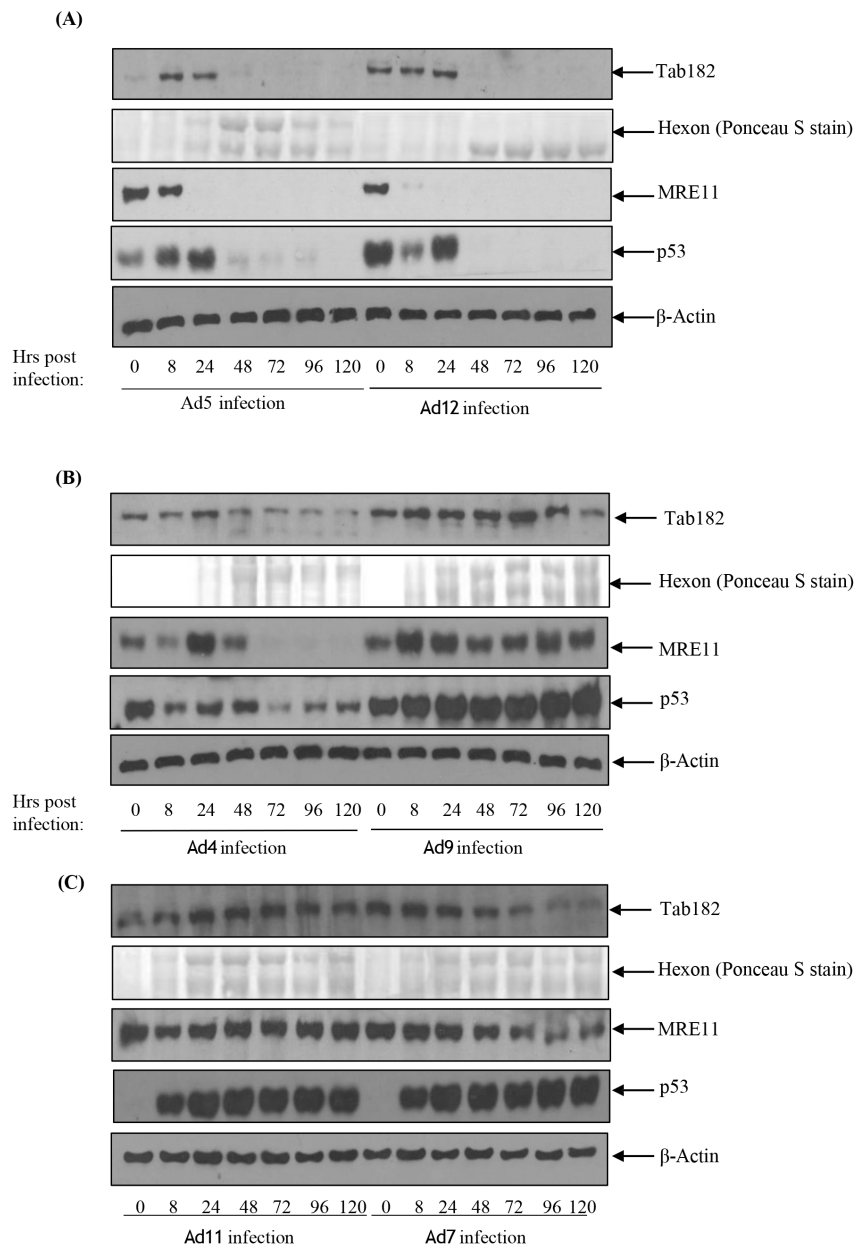


Figure 5

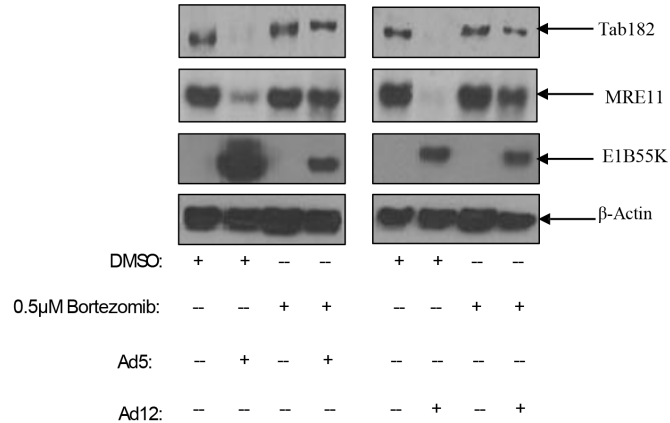


Figure 6

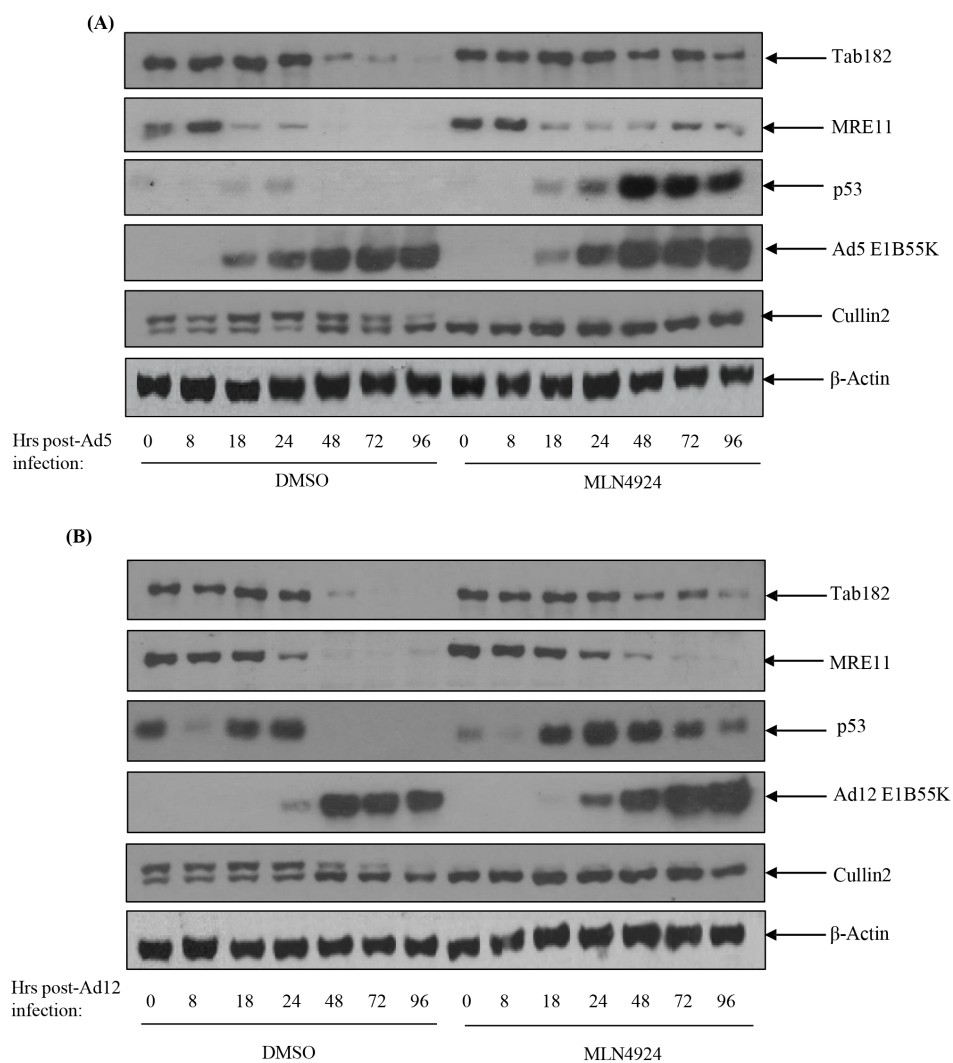


Figure 7

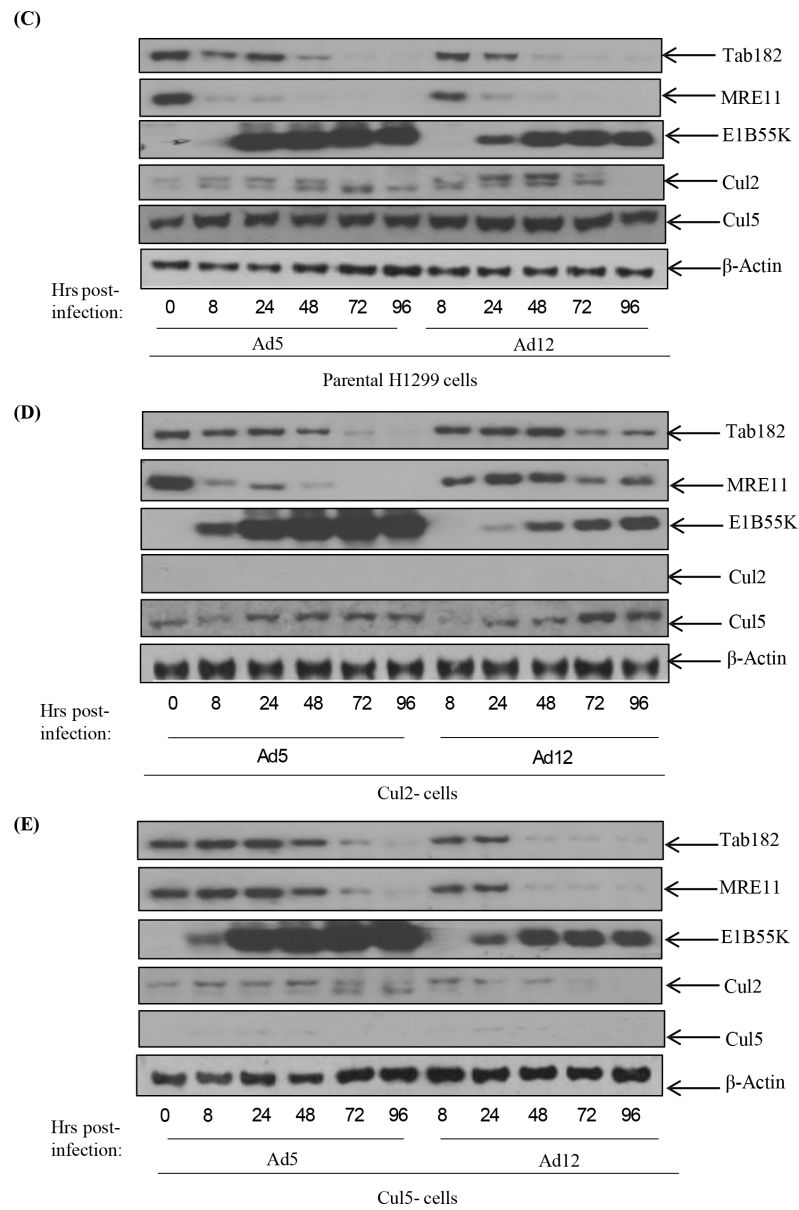


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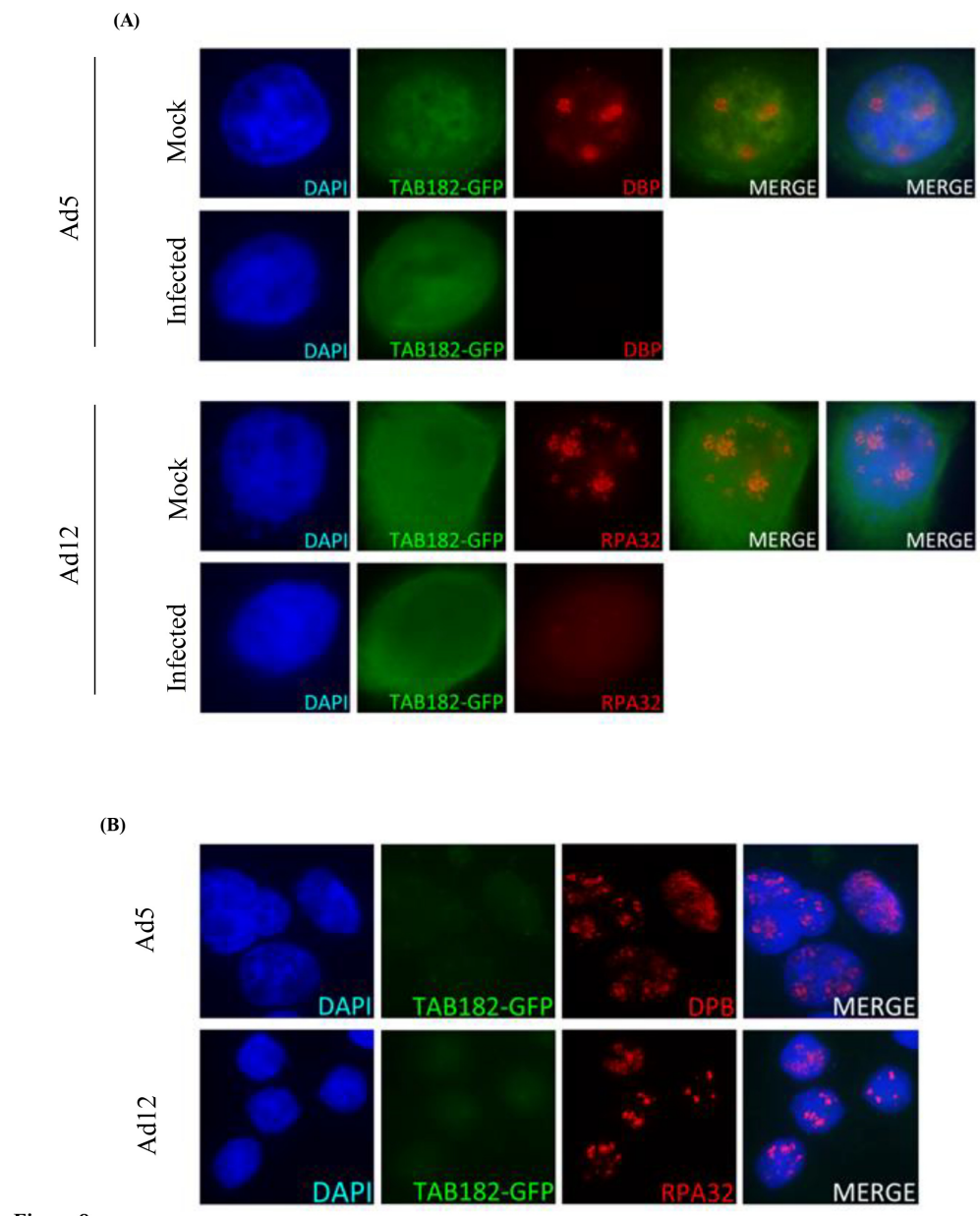


Figure 8

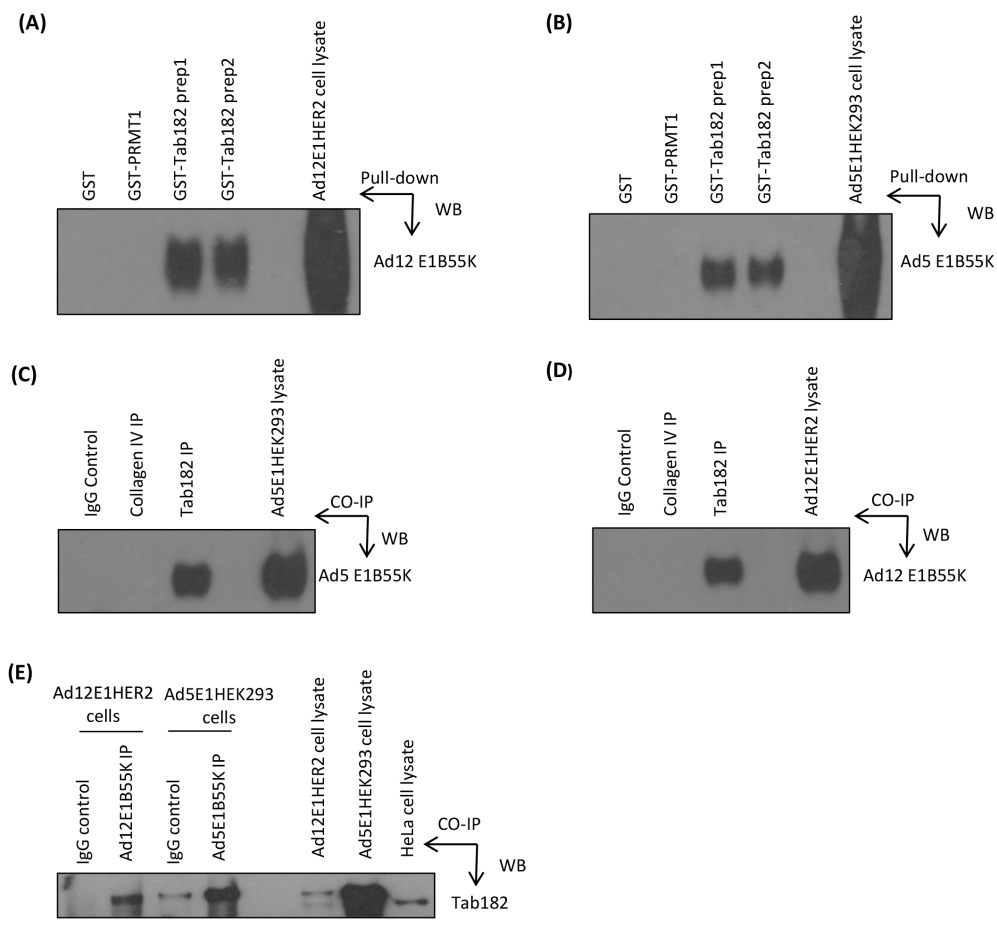


Figure 9

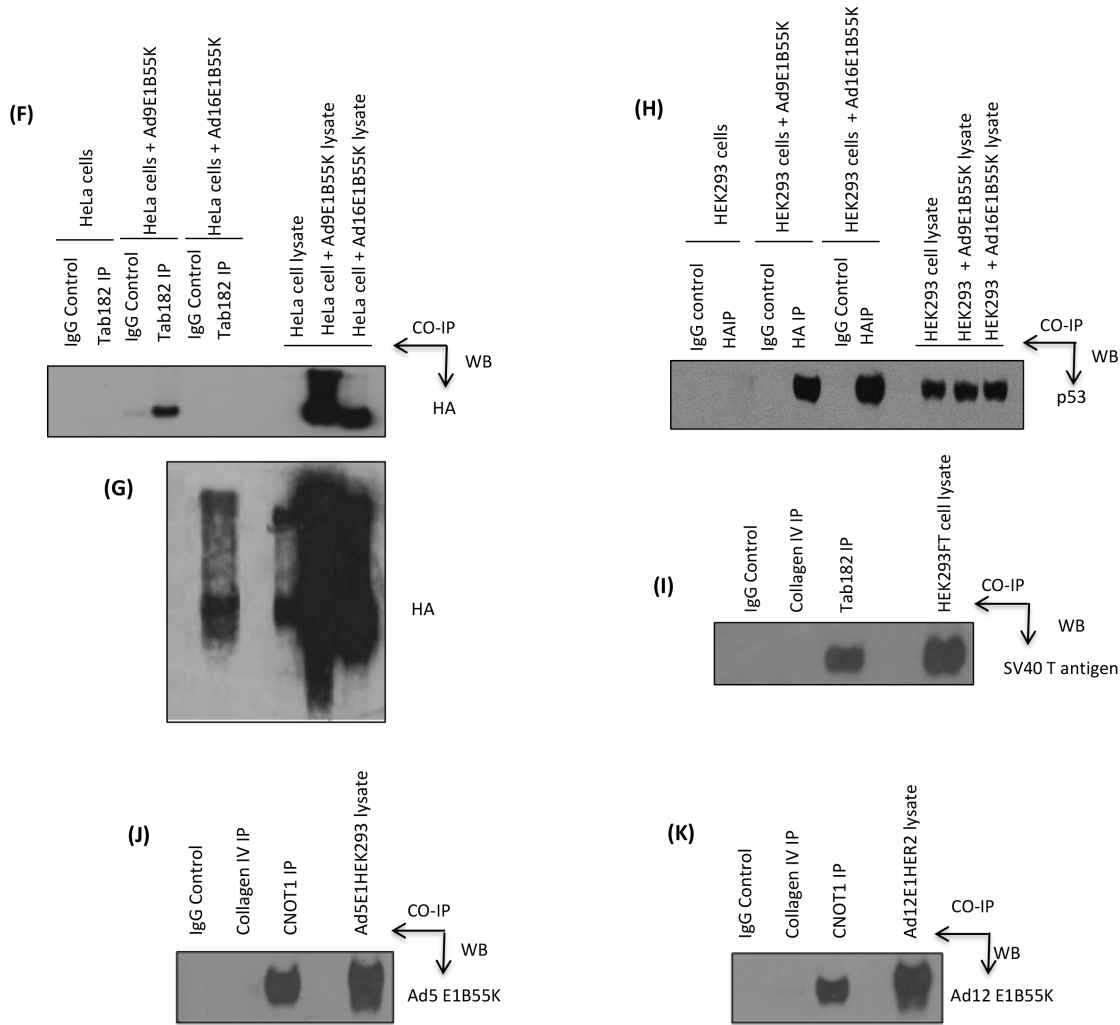


Figure 9

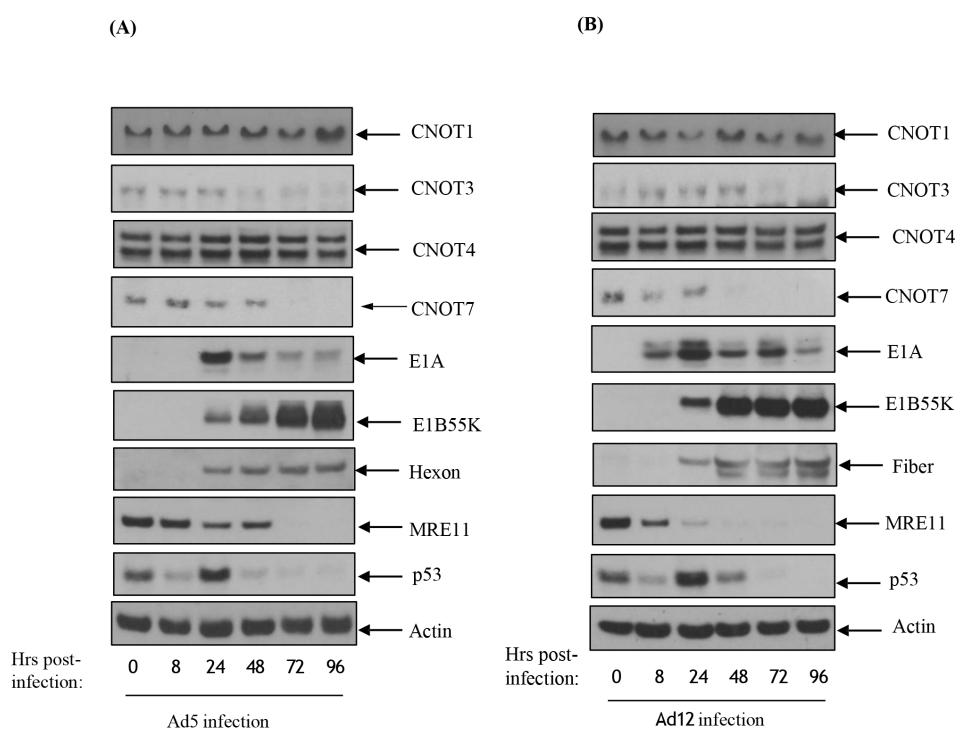


Figure 10

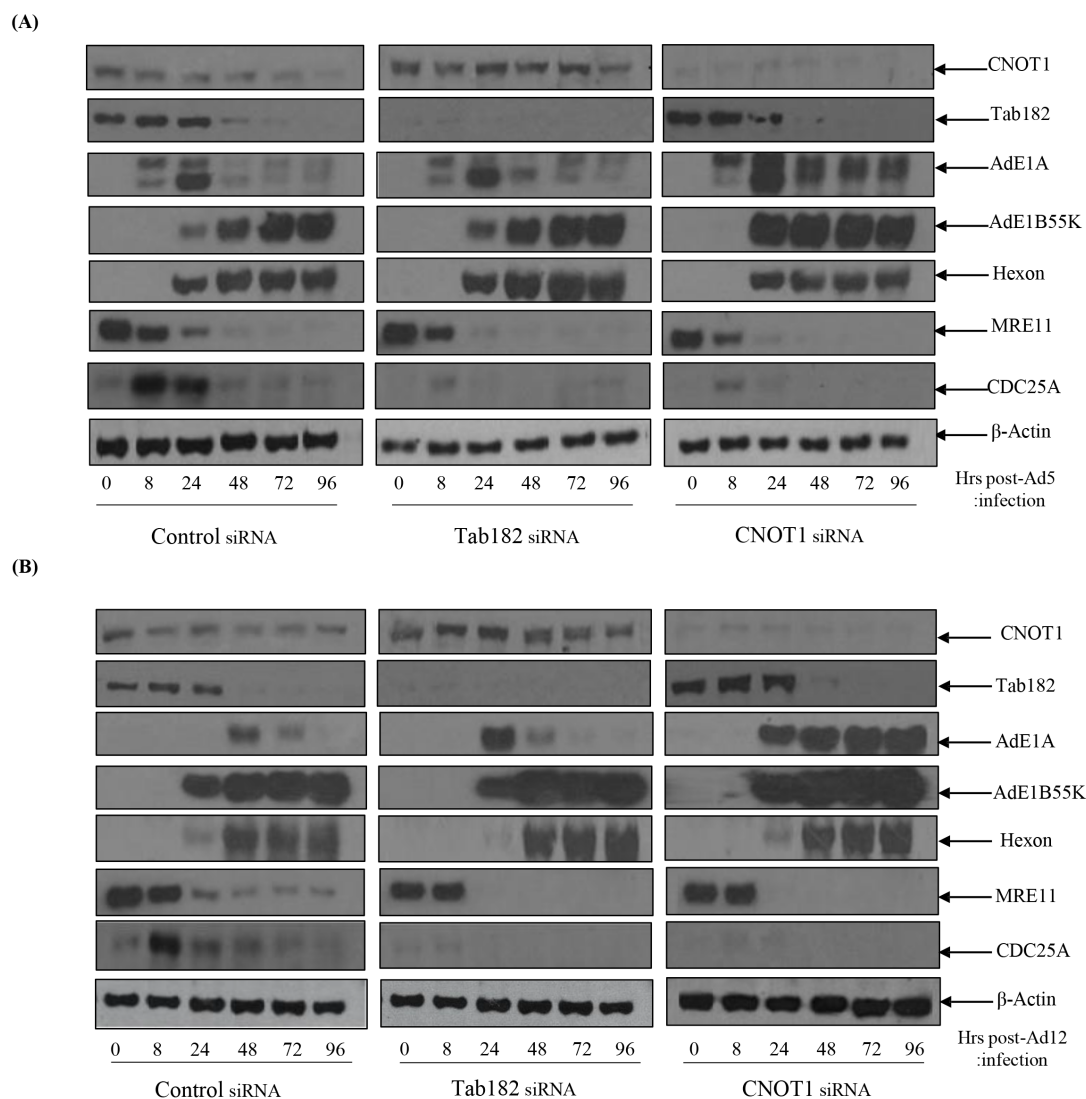


Figure 11

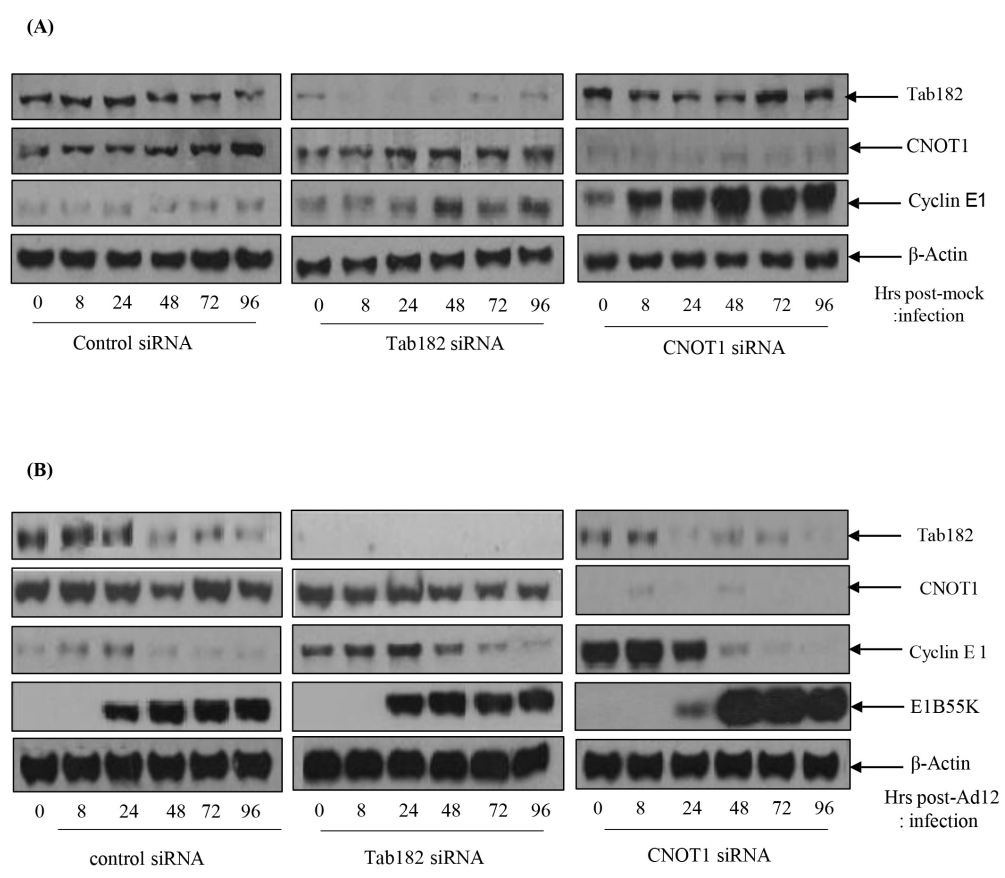
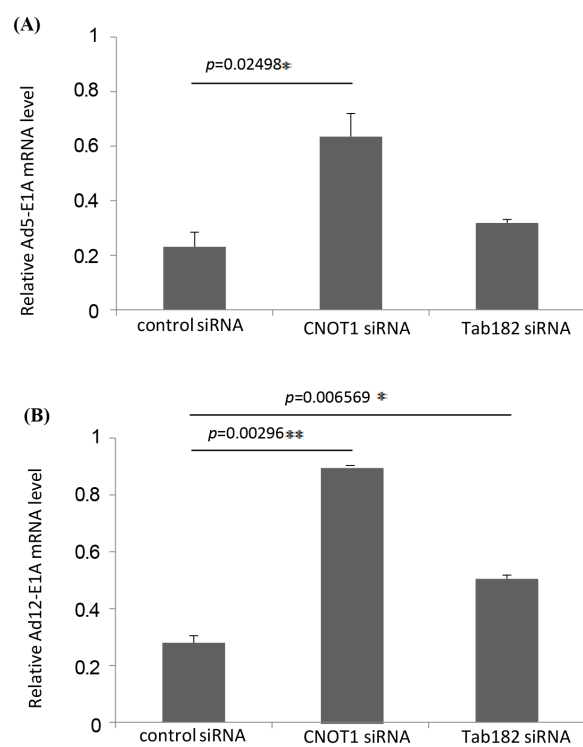
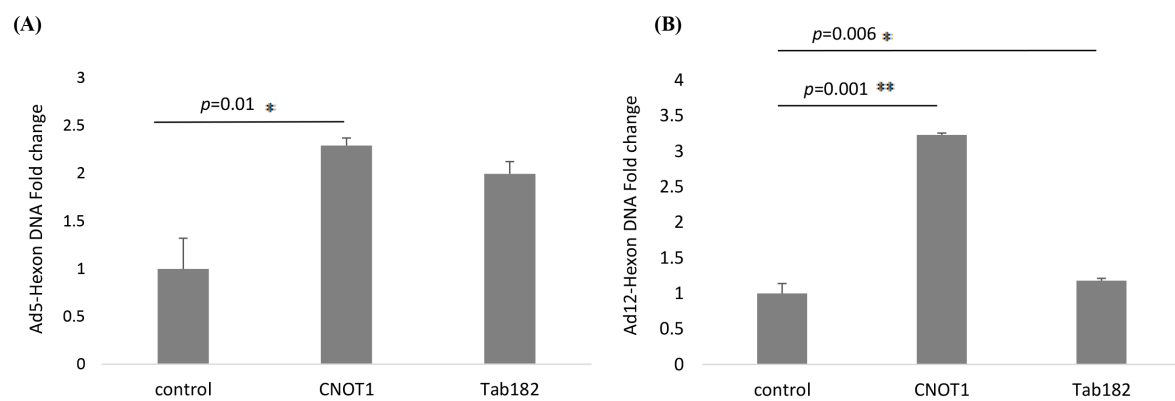


Figure 12

**Figure 13**

**Figure 14**